This is a "preproof" accepted article for *Journal of Clinical and Translational Science*. This version may be subject to change during the production process. 10.1017/cts.2025.56

How do study participants want to be informed about study results: findings from a malaria trial in Cambodia, Ethiopia, Pakistan and Indonesia

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Abstract

Background. Researchers acknowledge the need to share study results with the patients and their communities, but this is not done consistently due to a plethora of barriers, including a paucity of data to guide best practice approaches in different populations.

Methods. This study was nested within a large multi-centre randomized controlled trial of antimalaria treatment. Data on dissemination preferences were collected at the third month follow-up visit using a short questionnaire. Data were analysed using descriptive statistics and subsequently fed into an iterative process with key stakeholders, to develop suitable strategies for result dissemination.

Results. A total of 960 patients were enrolled in the trial, of whom 84.0% participated in the nested survey. A total of 601 (74.6%) participants indicated interest in receiving trial results. There was significant heterogeneity by study country, with 33.3% (58/174) of patients indicating being interested in Cambodia, 100% (334/334) in Ethiopia, 97.7% (209/214) in Pakistan, but none (0/85) in Indonesia. The preferred method of dissemination varied by site, with community meetings favoured in Ethiopia (79.0%, 264/334) and individualised communication such as a letter (27.6%, 16/58) or phone calls (37.9%, 22/58) in Cambodia. Dissemination strategies were designed with key stakeholders and based on patient preferences but required adaptation to accommodate local logistical challenges.

Conclusion. The varying preferences observed across different sites underscore that a onesize-fits-all approach is inadequate. Strategies can be tailored to patient preference but require adaptation to accommodate logistical challenges.

Keywords: results dissemination; research participants; clinical trial, malaria, community engagement

Introduction

Community engagement is increasingly recognized as an integral element of ethical global health research ¹. Engaging community members in health research occurs on a spectrum, and their involvement as early as possible to set priorities for research and co-design relevant research is critical ^{1,2}. However, involvement of community members is often deferred until the start of the recruitment processes, in line with the instrumental utility of undertaking community engagement to commence the study and ensure patient participation ^{3,4}. Continued community engagement after the completion of research studies, particularly dissemination of aggregated non-individualised study results to research participants, is an ethical obligation ^{3,5} and regulatory requirement ^{6,7}. Some research funding bodies now recognise the importance of these activities and require integration of result dissemination activities into research proposals ⁸.

Disseminating trial findings can improve health literacy and decision-making among participants, improve general understanding of research and encourage participation in future research ^{9,10}. Furthermore, increased transparency and trust in medical research, can lead to better satisfaction among participants¹¹⁻¹³. While most researchers acknowledge the need to share aggregated study results with the patients and their communities, this is not practised consistently ^{14,15}.

A survey among malaria researchers showed that although more than 80% appreciated the importance of sharing results with the trial participants, only 25% accomplished this in their most recent trials ³. Key barriers to result dissemination in those settings include difficulty locating and reaching research participants after the end of the trial ¹⁶⁻¹⁸, low literacy levels among study participants ^{3,19}, limited advanced planning^{3,14,15}, logistical issues such as limited access to and availability of internet and phone, a lack of electricity and poor road conditions in resource-constrained settings, financial constraints ^{14,15}, a lack of institutional guidance on how to conduct dissemination activities ^{3,19,20} and ethical concerns, including concerns around confidentiality within small communities ²¹, and fear of misinterpretations and inflicting harm ^{14,15}.

There is a paucity of data to guide best practice for the dissemination of research findings in different study populations and settings. More commonly reported types of results sharing include written communication in letters or lay summaries^{22,23}, emails ²³, and information placed on websites ²⁴ or group presentations in the form of community meetings or

workshops ^{9,25,26}. The limited literature from low- and middle-income countries comes primarily from the African continent and focuses on experiences with community meetings as the main method for result sharing ^{16,25}. Most of the literature guiding the selection of methods as well as patient preferences is from high-income countries, and these methods are less likely to be relevant to participants from communities with structural barriers such as low health literacy, vulnerability, poverty, competing priorities, and access and interest to engage in learning about study findings.

Our multi-centred antimalarial trial in Cambodia, Ethiopia, Pakistan, and Indonesia was designed to compare the effectiveness, safety, cost-effectiveness, and feasibility of novel treatment options for patients with *Plasmodium vivax* malaria. As part of the trial, we conducted a nested study to assess trial participants' preference on result dissemination after study completion to inform result sharing strategies.

Methods

Study overview

The study was conducted in two stages. In the first stage (the 'survey'), data were collected on patients' preferences for dissemination. Following the analysis of these data, the second stage involved reviewing the findings of the survey and developing suitable dissemination strategies through an iterative process among the study team.

Study context and sites

Data for the survey were collected in the context of a multi-centre clinical trial to assess the safety and effectiveness of novel approaches to the treatment *P. vivax* with tafenoquine and primaquine (NCT04411836). In brief, adult patients presenting with uncomplicated *P. vivax* malaria meeting the eligibility criteria were randomized into one of three treatment arms. After a standardised informed consent process each patient was treated with a blood-stage treatment plus either low dose primaquine (total dose 3.5mg/kg) unsupervised over 14 days, high dose primaquine (total dose 7 mg/kg) unsupervised over 7 days, or a single dose of tafenoquine (300mg). Patients were then followed weekly until day 42 and then monthly for 6 months. Patients' recruitment occurred at seven study sites in four countries: Cambodia, Ethiopia, Pakistan, and Indonesia (Figure 1). Training on data collection and study conduct

was standardised across study sites. Background details on the study countries, malaria burden, their socio-economic structure and literacy are summarised in Table 1.

Cambodia

In Cambodia, patients were recruited at three sites: Phnom Kravanh referral hospital and Siem Pang and Chambak health centres. Phnom Kravanh referral hospital has a catchment population of more than 50,000 and is located in Pursat province, in western Cambodia, with the population concentrated around the Pursat river, and the western parts of the Cardamom mountains. Most communities in this area live in the forest fringe or nearby forested mountains. Although communities have access to health centres and larger health facilities, the distances between communities and these health centres can be far. Siem Pang health centre has a catchment population of 25,000 and is located within Stung Treng province in north-eastern Cambodia, along the Tonle Kong river, surrounded by forest reserves, and national parks with abundant forest fringes. A high proportion of ethnic minorities live in Siem Pang, with difficult access to health centres, hampered by road conditions, terrain, and long distances. Chambak health centre has a catchment population of 4,000 and is located within Phnom Sruoch, a district located within Kampong Speu province in central Cambodia. The district covers the largest part of the Kirirom National Park located in the eastern part of the Cardamom mountains. Most settlements are thus close to the forest or forest fringe. Although Chambak health centre is relatively accessible, access to larger health facilities specially among populations living in far flung communities near the forest is limited. The majority of the study team at the Cambodia sites was from within the communities where patients were recruited from. In addition, international staff was present at the study sites.

Ethiopia

The study site in Ethiopia was located at the research facility at the Arba Minch General Hospital, Arba Minch town, in the Gamo Zone of the South Ethiopia Region (SER). Arba Minch district is located 505km south of Ethiopia's capital, Addis Ababa, with a population of 320,000. The urban centre has a general hospital, a primary hospital, and two health centres. In addition there are a range of private facilities, at which approximately 20% of malaria patients seek care ²⁷. The study hospital receives patients from two health facilities located in Arba Minch town, Woze health centre and Dil Fana Primary Hospital. For urban patients, all health facilities are within walking distance or accessible by taxi or private car.

The entire study team at the Ethiopian site was from within the communities where patients are recruited from.

Pakistan

In Pakistan the study recruited patients at two sites: an urban site in Karachi and a more rural setting in Thatta. The Karachi site is at Khidmat-e-Alam Medical centre, Nazimabad, which is located in the densely populated central district of Karachi with an estimated population of nearly 3 million. This small health care facility serves a densely populated urban area, where residents face challenges accessing health services due to overburdened facilities. The site in Thatta was at the Thatta Civil Hospital, in Sindh province with an estimated catchment population of 979,817. This rural area is characterised by scattered housing and agricultural lands, with limited access to and longer travel distances for medical care. Some members of the study team at the Pakistan site were from within the communities where patients are recruited from, other team members commuted daily to the study site from Karachi.

Indonesia

The Indonesian study site was located at the primary health care centre in Batubara in Batubara Regency in North Sumatera province. Batubara Regency is located at the eastern shoreline of North Sumatera, with Limapuluh as its administrative seat. It covers an area of 887 km² with a population of 430,533. The health centre serves a catchment population of nearly 34,000 people and is the primary public health facility. Approximately 45% of the population seek treatment at private facilities (personal communication head of district health office Batubara). The study team was external to the communities where patients were recruited from and worked there for the duration of the study.

Data collection

At the scheduled follow-up visit three months after treatment, a short survey questionnaire was used to assess patients' preferences for study result dissemination. The questionnaire was designed based on earlier work exploring current practise of result dissemination among malaria researchers ³ and discussion among the site investigators. Patients were asked four close-ended questions by the study nurses on the following topics: i) interest in knowing the results of the trial in which they were participating (primary outcome), and if so ii) their motivation for this, iii) their preference for methods used for sharing results and iv) suitable content for the dissemination (multiple options could be selected). Following translation and

pre-testing of the questionnaire among the investigators and their teams, minor adjustments were made, mostly to reduce ambiguity (Text S1). Demographic characteristics from study participants were collected as part of the data collection for the trial at enrolment into the study.

Data analysis

Basic descriptive statistics were used for each question and analysis conducted overall and per country. A multiple linear regression analysis was used to identify predictors for the primary outcome. All statistical analyses were performed using Stata version v17.0 (StataCorp, US).

Development of dissemination strategies

Strategies for result dissemination were developed for each study location separately based on the results on patient preferences through an iterative process between the respective site team and the trial coordinating team. This iterative process included a virtual presentation of the country specific findings to the site study team, followed by a discussion on how results could be translated into a strategy considering the feasibility of the preferences. Based on suggestions and agreements in the virtual meeting, a written summary strategy was drafted and shared with study teams for feedback and further discussion via email. Where necessary additional virtual meetings were conducted until agreement was reached among all team members.

Ethics. The study was approved by the Human Research Ethics Committee (HREC) of Northern Territory Health and Menzies School of Health Research (#20-3694) and country-specific institutional, national and regulatory authorities (Table S1). All patients provided informed consent.

Results

Study population

A total of 960 adult patients were enrolled into the clinical trial, 220 in Cambodia, 350 in Ethiopia, 240 in Pakistan, and 150 in Indonesia. Data on dissemination preferences was obtained from 806 (84.0%) of the trial participants, of whom 174 (79.1%) were recruited in Cambodia, 334 (95.4%) in Ethiopia, 214 (89.2%) in Pakistan, and 84 (56.0%) in Indonesia. Non-participation in the survey only occurred when the scheduled follow-up visit at month

three post treatment was missed. The demographic characteristics of patients who participated in the survey were similar to those for whom no data was collected (Table 2, Table S).

Interest in learning about study results

Overall, 74.6% (601/806) of participants indicated they were interested in learning about the study result. This was different by study country with 33.3% (58/174) of participants in Cambodia being interested in receiving aggregated study results, compared to 100% (334/334) in Ethiopia, 97.7% (209/214) in Pakistan, and 0% (0/85) in Indonesia (p<0.005). A total of 82.3% (210/255) female patients compared to 71.1% (391/551) male patients indicated interest in learning about study results (p=0.001). Interest across age groups was distributed as follows: 100% (15/15) among the 16–18-year-olds, 79.8% (400/501) among those 18 to 30 years, 68.6% (192/280) among patients 30 to 60 years and 90% (9/10) among those over 60 years old (p<0.001).

In the multivariable analysis the differences between sites remained significant (p<0.001) after controlling for age and sex.

Motivation for hearing about study results

In total 45.6% (274) of the 601 patients who expressed an interest in hearing about the study results indicated that the main reason was to understand the benefit of the study to the community. A further 16.3% (98) patients indicated they were interested to understand the study in an accessible and easy way, and 38.1 % (229) patients indicated that it represented an acknowledgement of their contribution to the study. Motivation for wanting to hear results differed by country, with 76.6% (160/209) of patients in Pakistan indicating acknowledgement of their contribution as the most important reason, while understanding the benefit of the study to the community was selected by 71.9% (240/334) patients in Ethiopia (Table 3).

Preference for dissemination methods

Participant preference for methods by which study results are communicated varied by country. A large majority of patients in Pakistan (99.0%; 207/209) felt that it was important to have summaries in their own language, while this was only reported by 38.3% (128/334) in Ethiopia. Cambodian patients preferred receiving a letter (27.6%; 16/58) or a phone call from someone explaining the study results (37.9%; 22/58), while Ethiopian patients

overwhelmingly preferred a community meeting at the clinic or health centre (79.0%; 264/334). In Pakistan, a high preference was indicated for having the results published on a website (60.8%; 127/209), a personal phone call to explain results (76.1%: 159/209) or a community meeting (92.3%; 139/209) (Table 4).

In Cambodia and Ethiopia, most patients selected three or less options for methods of dissemination (57/58, 98.3% and 329/334, 98.5%), whereas in Pakistan 66.0% (138/209) selected more than 3 options (Figure S1-3).

Preference for dissemination content

Almost all patients in Pakistan (94.3%, 197/209) and Ethiopia (93.7%; 313/334) felt that it was important to include information about the purpose of the study in the dissemination summary. Whereas patients in Pakistan felt that it was important to include information about medical treatment (84.7%; 117/209) and scientific (75.6%; 158/209) advances based on the study's results. Patients in Ethiopia were interested to hear about how the study was conducted (39.5%, 132/334) and the potential implications of the results to change treatment policy (30.2%; 101/334). Cambodian patients had limited interest to hear about general scientific advances (1.7%; 1/58), potential policy change (0%; 0/58) or new research based on the results (0%; 0/58). Overall patients felt it was more important to include positive findings (72.1%; 333/601) compared to negative (11.0%; 66/601) or neutral results (7.3%; 44/601) (Table 5).

Dissemination strategies

Separate dissemination strategies for Cambodia, Ethiopia, and Pakistan were developed based on the survey results to inform the target audience, the method of delivery and the content. No strategy was developed for the Indonesian site, given the lack of local interest. Although 23.2% (126/601) of participants indicated that they want to hear about individual results, this was not within the scope of our dissemination efforts. Draft strategies are summarized in Table 6.

Discussion

Our study highlights marked heterogeneity between study countries in the interest of hearing about the results of the clinical trial that they had been enrolled into, preferred methods of dissemination, and the content to be included in dissemination activities. None of the Indonesian patients expressed an interest in hearing about the study results, while all of the Ethiopian patients indicated interest. In Ethiopia the preferred method for dissemination was a community meeting, whereas in Cambodia patients preferred more individualised communication. In Pakistan patients had strong preference for community meetings, individual communication as well as web-specific distribution. While most patients wanted to hear about study results in their own language and wanted to be reminded about the purpose of the study, there were large differences in the degree of interest about the impact on medical or scientific advances and the implications for policy.

Our study underscores the importance of contextualizing dissemination strategies according to social, cultural and research literacy related characteristics of the population rather than adopting a one-size-fits-all approach. The results also indicate the potential added benefit to conduct formative research alongside clinical trials so that community preferences can be explored, and community engagement strategies tailored to local populations ^{28,29}. Previous studies have demonstrated that formative research is deemed to be critical for optimal design and implementation of interventions and programs ³⁰⁻³³. However, in practice these processes are constrained by the available resources and how they can be used, which in turn affects the degree of involvement of research participants ². Community engagement usually occurs on a wide spectrum of activities from simply asking about preferences to incorporating community opinions at study inception through co-design ³⁴⁻³⁶. While our survey focused primarily on exploring participants preferences for post-trial engagement, these engagement activities (exploring participants' preferences) even if they are subtle, can demonstrate respect, and forge providence for future research, sustaining trust and relationships ³⁶⁻³⁸.

The ethical obligation to disseminate research findings to participants, can conflict with respecting the agency and autonomy of patients, particularly if they prefer not to be informed about the trial results ³⁹. None of the Indonesian study participants indicated interest in hearing about the trial results and therefore no dissemination strategy was developed for this study population. While further qualitative research is required to fully understand motivations and drivers, a potential explanation for this finding based on internal team discussions might be the fact that the study team was external to their community which may have fostered a sense of difference: 'us' versus 'them' thus deterring community members from further engagement on results dissemination. Other generic barriers potentially include socio-economic constraints that restrict participants time, and opportunity costs incurred whilst engaging in results-dissemination efforts. These factors may also have contributed to the relatively large number of participants who were lost to follow up at the Indonesian site.

However, it remains unclear how much of a role this played compared to other sites where patients are coming from similarly low socio-economic backgrounds.

Preferences of engagement activities can vary widely based on cultural, educational, and infrastructural factors ^{9,19,29,40}. For example, the strong preference for community meetings in Ethiopia and Pakistan is consistent with studies indicating that face-to-face communication is often more effective in low-resource settings where digital literacy and internet access may be limited ^{41,42}. The results are in line with previous practise at the study site in Ethiopia ²⁵, potentially indicating that participants selected choices they are more familiar with. While there was a clear preference for community meetings in Ethiopia, in Pakistan a preference for individual communication was also stated, which aligned more with Cambodian patients. The preference for personalized communication in Cambodia likely reflects participants desire not to be engaged frequently, owing to their involvement in agricultural work coupled with barriers to reaching the health centre ^{43,44}. Sometimes, community members may feel participation in research and engagement itself as a burden ¹. Cambodian participants have been reported to exhibit social tendencies of conformism, quietness, and hierarchical etiquette with researchers and a reluctance to 'losing face' ⁴⁵. In response to these social and cultural characteristics and to bridge the differences between researchers and community members, in Siem Pang, a youth advisory group on health and research engagement was recently established ^{36,37}.

Patients at the Pakistani study sites were the only ones indicating a high preference for digital communication, including a webpage or digital messaging services. This preference was evident in both urban and rural settings. Given that the large majority of patients were recruited in the rural site, no meaningful comparison between sites was possible (data not shown). Given the relative low literacy rate in Pakistan coupled with digital inequality ⁴⁶ these results are surprising, requiring further exploration. Digital communication allows reviewing information at convenient times for the individual and reducing additional opportunity costs for travel and in person meetings, which might be relevant in this patient cohort.

Our findings also revealed heterogeneity in preferences about the content of the dissemination. While there was a high interest to learn more about implications for the community in Ethiopia in line with the preference for community-based delivery methods,

there was a lack of interest for policy implication in Cambodia, which might be explained as a function of a research saturated setting.

The initial design of our dissemination strategies was based on patient preferences; however, adaptations were needed to accommodate the realities of what investigator teams considered logistically and financially feasible. For example, individual phone calls to each participant were considered impractical by the study team in Pakistan. This highlights that patient preferences do not necessarily align with what investigators perceive as being feasible.

A substantial number of patients indicated they were interested in learning about individual test results arising from the trial. Although some participants may conflate the difference between sharing of aggregated and individual results, the impact for participants is likely to be similar ⁴⁷. There are however a number of considerations that are distinctly different between sharing aggregated study results versus individual test results including concerns around confidentiality, interpretation of results, and implications for further care ⁴⁸. Most of the research on returning individual test results to study participants comes from genomic research where interpretation of results remains challenging ⁴⁹. However, studies indicate that some participants prefer to receive individual results even if they have no clinical significance ⁵⁰. Our preliminary qualitative research conducted in Cambodia (unpublished data) suggests that communication on individual results during the follow up is important to patients and could affect perceptions on dissemination of overall study results and engagement in future trials.

Our study has several limitations. Firstly, the questionnaire was developed based on previous work assessing current practise of result dissemination among malaria researchers ³, but was not co-designed through formative qualitative work with patients. Therefore, the selection of choices both for methods as well as content of dissemination may not have included the entire breadth of possibilities relevant for participants. However, none of the participants used the 'other' option that was designed to capture additional concepts. Second, no data on socio-economic, ethnic or educational background was collected. This would have allowed for a more granular analysis of our findings for different populations. Third, in line with the limitations of quantitative surveys the data does not provide explanations as to the reasons behind the preferences and more qualitative research would be needed to explore this in depth. Fourth, although training was standardised across study sites, we cannot exclude that answers to questions were not influenced by social desirability bias, or by the way study

teams asked questions differently depending on their preconceived ideas. This includes potential differences in the extent of information that was provided as part of the consenting process. Fifth, patients recruited into the trial and surveyed do not necessarily reflect the overall population and thus their preferences may not be generalizable for the larger population, this includes potential differences by gender. The majority or participants in the trial were male and this bias towards male patients was particularly strong in the Cambodian site. This reflects the epidemiology across the Greater Mekong subregion where malaria is predominantly a disease of young males with increased occupational risks (e.g. forest goers) to get infected ⁵¹. Lastly, at the time of writing the results of the trial were not yet available, therefore, the developed dissemination strategies include only high-level guidance on content. In addition, no evaluation of the impact and feasibility of the different strategies has been conducted.

In conclusion, our study highlights the critical need for tailored dissemination strategies in global health research. The varying preferences observed across different countries underscore that a one-size-fits-all approach is inadequate. Researchers must consider patient preferences and context when planning dissemination activities.

Author contribution. KT and HU conceptualised the study, KT and RNP acquired funding, SAB, FJ, NH, TTT, WA, Y, HM, and BA collected data, SAB, FJ, BA and KT conducted data analysis, TSD, NG, MAB, RT, APP, HM, GL, LS, LvS, HU, BA and KT supervised the study conduct, SAB, FJ and KT developed the first draft, MBV, NH, HU, BA and RNP contributed to the writing, all authors reviewed and approved the final manuscript.

Acknowledgements. We would like to thank all patients and their communities.

Competing interests. The authors declare none.

Funding statement. This study was supported through a grant from the National Health and Medical Research Council (KT, grant number 1182950) and from the Bill and Melinda Gates Foundation (RNP, INV-024389). KT is a CSL Centenary Fellow, and RNP is supported by an NHMRC Investigator Grant (2008501). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Table 1: Background on study countries

	Cambodia	Ethiopia	Pakistan	Indonesia
GDP per capita (US\$)*	1,759.6	1,027.6	1,588.9	4,788.0
Life expectancy at birth in years (2021)*	70	65	66	68
Literacy rate, adults total (% of people ages 15 and above) *	84% (2022)	52% (2017)	58% (2019)	96% (2020)
Mobilephonesubscriptionper 100people (2022)*	116	56	82	115
Fixedbroadbandsubscriptionsper100 people (2022)*	3.04	0.46	1.33	4.88
Social Media use in percent of population (2024) 1	68.4%	5.5%	29.5%	49.9%
Occupation and livelihood at study sites	Primarily agriculture. Community members are busy during rainy season planting rice and vegetables and may find difficult to travel during the season because of road conditions.	Mainly government, private, NGOs or self- employed. A smaller number of people in agriculture.	InurbanInruralKarachi,mostThatta,peoplearemanyengagedincommunitgovernment,y membersprivatesectorarejobs,NGOs,engaged in	Agriculture and fishery are primary occupations. A smaller number of people work as government employee or self- employed

			employed, with and related	
			a smaller activities	
			number	
			involved in	
			agriculture.	
Malaria disease	Malaria transmission is	Malaria occurs year-	Malaria occurs year-round,	Malaria occurs year-round
burden at study sites	heterogeneous. In 2022	round, mainly	with higher transmission rates	with higher transmission
	there was an outbreak of	following the rainy	following the rainy seasons.	during rainy season.
	falciparum malaria in	seasons.	In urban Karachi, malaria	Batubara is malaria
	Kravanh ⁵² . In between	The peak seasons for	prevalence is relatively low	moderate endemicity with
	October-December 2023,	malaria transmission in	(5 cases per 1000 population	1.49 API, and one of the
	there were a total of 276	the country is from	yearly) due to better	regencies that contributed
	cases (1% P. falciparum and	September–December.	healthcare infrastructure and	to highest malaria cases in
	mixed cases, and 94% P.	Over 80% of the	urban living conditions. In	North Sumatera province.
	vivax cases), a 70%	malaria burden in	rural Thatta, malaria remains	Ratio between P. vivax
	decrease compared to the	Ethiopia is among	a significant health issue,	and P. falciparum is 70:
	same time period in 2022 53 .	adults and children who	with higher prevalence rates	30 (provincial malaria
	Almost half of the malaria	are at least five years of	(25 cases / 1000 population)	data).
	cases between January to	age. The	attributed to agricultural	
	July 2024 (of total malaria	The Southern Nations,	practices and limited access	
	in Cambodia) were reported	Nationalities, and	to healthcare facilities ⁵⁷ .	
	from Stung Treng that hosts	Peoples' Region		
	Siem Pang district ⁵⁴ . Siem	(SNNPR) is		
	Pang had the highest	categorised as moderate		
	malaria cases (n=70)	to high disease burden		
	between January to July	with API between >10-		
	2024, and almost all of them	50 for areas categorised		
	were vivax malaria. Most of	as moderate and >50 for		

these vivax malaria cases	those categorised as	
were found among soldiers,	high ⁵⁵ . There were	
migrants, and ethnic	415,197 confirmed	
minorities. Phnom Sruoch	malaria cases reported	
that host Chambak health	from SNNPR in 2022,	
centre had high prevalence	and 126,029 (30%)	
of vivax malaria in 2022,	were attributed to P.	
but the prevalence since	vivax ⁵⁶	
then has declined with only		
three cases of malaria		
between January to July		
2024 in an entire district.		

*based on https://data.worldbank.org/indicator

lfrom https://datareportal.com/reports/digital-2024.

		Cambodia	Ethiopia	Pakistan	Indonesia	Total
		N=174	N=334	N=214	N=84	N=806
Sex	Male	155 (89.1%)	188 (56.3%)	155 (72.4%)	53 (63.1%)	551 (68.4%)
	Female	19 (10.9%)	146 (43.7%)	59 (27.6%)	31 (36.9%)	255 (31.6%)
Age in	16-<18	0 (0%)	0 (0%)	0 (0%)	15 (17.9%)	17 (2.3%)
years*	18-30	113 (64.9%)	256 (77.0%)	100 (46.7%)	31 (36.9%)	462 (62.6%)
	31-60	61 (35.1%)	76 (22.8%)	106 (49.5%)	37 (44.1%)	251 (34.0%)
	>60	0 (0%)	1 (0.3%)	8 (3.7%)	1 (1.2%)	8 (1.1%)

Table 2: Demographic characteristics of study participants by study country

*The study site in Indonesia allowed recruitment of patients ≥ 16 years, while the other study sites only recruited patients ≥ 18 years

Table 3:	Reason	for	dissem	ination	bv	study	country
1 4010 01	neubon	101	4100011	mation	$\mathcal{O}_{\mathcal{J}}$	Staaj	country

	Cambodia	Ethiopia	Pakistan	TOTAL
	N=58	N=334	N=209	N=601
To understand the benefit	24 (41.4%)	240 (71.9%)	10 (4.8%)	274
of the study to the				(45.6%)
community				
Acknowledgement of my	17 (29.3%)	52 (15.6%)	160 (76.6%)	229
contribution to the study				(38.1%)
To understand the study in	17 (29.3%)	42 (12.6%)	39 (18.7%)	98 (16.3%)
an accessible and easy way				

 Table 4: Preference for dissemination methods by study country

	Cambodia	Ethiopia	Pakistan	TOTAL
	N=58	N=334	N=209	N=601
I prefer a meeting at the	5 (8.6%)	264 (79.0%)	139 (92.3%)	462 (76.9%)
Clinic/ health center				
together with other trial				
participants and someone				
explain the results				
I prefer a summary in my	45 (77.6%)	128 (38.3%)	207 (99.0%)	380 (63.2%)
language				
I prefer someone to call	22 (37.9%)	7 (2.1%)	159 (76.1%)	188 (31.3%)
me and explain the study				
result				
Published on Website	0 (0%)	24 (7.2%)	127 (60.8%)	151 (25.1%)
Sent via Text/ WhatsApp/	9 (15.5%)	20 (6.0%)	43 (20.6%)	72 (12.0%)
Messenger				
I prefer someone to come	3 (5.17%)	19 (5.7%)	33 (15.8%)	55 (9.2%)
to my house and explain				
the study results in person				
Via letter	16 (27.6%)	1 (0.3%)	10 (4.8%)	27 (4.5%)
Via email	0 (0%)	14 (4.2%)	2 (1.0%)	16 (2.7%)
Published on Twitter/X	0 (0%)	7 (2.1%)	5 (2.4%)	12 (2.0%)

Table 5: Preference for content of dissemination by study count

	Cambodia	Ethiopia	Pakistan	TOTAL
	N=58	N=334	N=209	N=601
Purpose of the study	32 (55.2%)	313 (93.7%)	197 (94.3%)	542 (90.2%)
Good results	33 (56.9%)	214 (64.1%)	186 (89.0%)	333 (72.1%)
General scientific	1 (1.7%)	57 (17.1%)	158 (75.6%)	216 (35.9%)
advances based on the				
study's results				
Medical treatment	8 (13.8%)	67 (20.1%)	177 (84.7%)	152 (31.9%)
advances based on the				
study's results				
How the study was	4 (6.9%)	132 (39.5%)	31 (14.8%)	176 (27.8%)
conducted				
Potential policy	0 (0%)	101 (30.2%)	39 (18.7%)	140 (23.3%)
changes based on the				
study's results				
Potential new research	0 (0%)	84 (25.2%)	47 (22.5%)	131 (21.8%)
based on the study's				
results				
Results that are	18 (31.0%)	47 (14.1%)	61 (29.2%)	126 (21.0%)
specific to me				
Bad results	8 (13.8%)	29 (8.7%)	29 (13.9%)	66 (11.0%)
Neutral results	3 (5.2%)	6 (1.8%)	35 (16.8%)	44 (7.3%)

 Table 6: Draft dissemination strategies.

	Cambodia	Ethiopia	Pakistan
Target	All patients who indicated they are interested	All recruited patients	All recruited patients
Population	in hearing about the study results		
Approach	Patients will be selected from the study database based on their individual identifiers	Patients will be contacted using the contact information collected in the	A two-pronged approach will be used: firstly, patients will be contacted using
	and contact information will be derived from	patient information log and invited to a community meeting at the study centre	the contact information collected in the patient information log and invited to a
	was used to support patient follow up. If	Given the large number of patients	community meeting at the study centre.
	phone numbers are available and still	multiple such meetings might be	Given the large number of patients
	functioning, patients will be contacted by	conducted. In line with previous similar	multiple such meetings might be
	senior staff members and asked if they want	meetings at the study site , the site	conducted. Given the importance of
	to hear about study results on this phone can.	investigator will provide a verbal	acknowledgement of participant
	All option to arrange for a call at a fater time point will be provided. If patients agree a	answer session	as 'thank you' meetings as opposed to
	short pre-written summary will be presented		focusing on information sharing.
	verbally. Participants will also be offered		Secondly, an information package will be
	option to receive the written summary in		prepared to be shared on a webpage and
	addition or as an alternative to the phone		patient will be informed about this in
	conversation. This will be sent via		conjunction with the invite to the
	WhatsApp/ messenger.		community meeting. The online package
			will include written information as well
			as a video. This will be after publication
			of study results.
Additional	Given the history of the site conducting	In line with previous experience ^{25,58} the	The investigator team opted against

measures	community meetings to disseminate results	community meetings will be	providing information at individual bases
	⁵⁸ , participants will also be offered an option	accompanied by refreshments	given the logistical difficulties and time
	to come to the health centre for an additional	supporting further informal	commitments this would require.
	meeting to discuss on the study findings.	conversations aimed at helping	However, if participants request
		participants gain a deeper	additional information, they will be
		understanding of the study's goal and	offered a one-on-one meeting.
		potentially become community	
		advocates for forthcoming activities	
		including the new studies.	
Content	Verbal summary as well as the written	The verbal communication during the	Both the verbal communication during
focus	summary will include information on the	meeting will include information on the	the meeting as well as the online
	purpose of the study, medical treatment	purpose of the study, how the study was	information will be framed to
	advances based on the study's results as well	conducted, the implications for medical	acknowledge participants participation
	as a short summary of how the study was	and scientific advances as well as	and thank them for their contribution.
	conducted.	implications for new research and	Information provided will include
		policy. A focus will be given for	information on the purpose of the study
		participants to appreciate the benefits of	as well as the implications for medical
		the study to the community.	and scientific advances. Online
			information will only be available after
			the study results are published.



Figure 1. Generated using QGIS Lima and finalized in Canva Pro. Country Shapefiles were obtained from the Humanitarian Data Exchange (<u>https://data.humdata.org/dataset/cod-ab-pak?</u>, <u>https://data.humdata.org/dataset/cod-ab-khm</u>, <u>https://data.humdata.org/dataset/cod-ab-eth</u>, <u>https://data.humdata.org/dataset/cod-ab-idn</u>) and are licensed under a Creative Commons Attribution 4.0 (CC-BY 4.0) International licence. The world map shape file was obtained from Opendatasoft (<u>https://public.opendatasoft.com/explore/dataset/world-administrative-boundaries/export/</u>), and license under an Open Government License v3.0.