Accelerating the elimination of *P. vivax* malaria in the Americas

About PAVE

The Partnership for Vivax Elimination (PAVE) led by PATH and Medicines for Malaria Venture (MMV) brings together National Malaria Programs (NMPs), researchers, funders, and other organizations to support countries as they strive to achieve their elimination goals through:

- Working with NMPs to identify optimal radical cure tools and strategies for their given contexts.
- Generating and sharing high-quality evidence on *Plasmodium vivax* (*P. vivax*) case management to inform policy decisions and implementation by national governments.
- Advancing the development and availability of quality-assured medicines and diagnostics for *P. vivax*.

Eliminating *P. vivax* malaria

Eliminating malaria is an ambitious goal, and relapsing *P. vivax* malaria presents a major obstacle to its achievement. In countries approaching elimination, the decline in *P. vivax* burden often lags the decline of *P. falciparum*. The *P. vivax* parasite has a dormant liver stage—the ‘hypnozoite’—that can be activated weeks, months, or years after the initial infection, making it difficult to eliminate. Diagnosis, and radical cure treatment requires coordinated use of (1) a malaria blood-stage diagnostic (either a rapid diagnostic test or microscopy), (2) a glucose-6-phosphate dehydrogenase (G6PD) diagnostic test, (3) a blood-stage antimalarial, either chloroquine (CQ) or an artemisinin-based combination therapy (ACT) (based on national strategies), and (4) tafenoquine (TQ) or primaquine (PQ) to clear parasites from the liver and prevent relapse. TQ and PQ can both cause severe hemolysis in G6PD deficient patients. As a result, G6PD testing is mandatory before using TQ, as patients can only be treated if they have enzyme activity ≥70%. G6PD testing is also recommended before using PQ to guide appropriate use and dosing.

New Tools for Radical Cure

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Photo: Jamil Barton/ PATH
Quantitative test to measure G6PD activity
The STANDARD™ G6PD Test provides a quantitative measure of a patient's G6PD enzyme activity levels. The SD Biosensor handheld device provides a patient’s G6PD level and hemoglobin measurement in 2 minutes based on a finger-prick. As of August 2022, the test is registered/available in over 18 countries. In addition, the test received Australian Therapeutic Goods Administration (TGA) approval in 2021 and Global Fund Expert Review Panel for Diagnostics approval until July 2023.

Tafenoquine—a single-dose relapse prevention treatment
TQ is a single-dose liver-stage treatment that provides radical cure for *P. vivax* malaria when used in combination with CQ. The Australian TGA and United States Food and Drug Administration approved TQ in 2018 for patients aged 16 years and over. It is now registered for the same population in Brazil, Colombia, Peru, and Thailand. In March 2022, the Australian TGA approved the use of TQ for children aged 2 years and above. The approval includes a novel 50 mg tablet that can be dispersed in water to facilitate use in children.

Pediatric primaquine for relapse prevention
PAVE is supporting Fosun Pharma to develop a 2.5 mg and 5 mg dispersible PQ tablet addressing the supply gap in pediatric treatment for relapsing malaria. If approved by WHO prequalification, it will be the first quality-assured pediatric PQ formulation available.

Generating evidence on *P. vivax* case management
In support of ministries of health, and led by in country research partners, PAVE is conducting feasibility studies to investigate the best use of *P. vivax* treatments and diagnostics to support malaria elimination. For more information visit the *P. vivax* information hub.

PAVE collates evidence to support national policy decisions of malaria-endemic countries. Country landscape reports have been developed or are underway for most *P. vivax* endemic countries, including Guatemala, Honduras and Panama. Visit the study database for an overview of vivax research studies; and the G6PD Operational Research Community of Practice site for practical information regarding the use of the G6PD test.

Progress in the Americas
- In Brazil, the Tafenoquine Roll-out STudy (TRuST) has been completed and results are expected by Q4 2022, alongside health economic assessments (HEA) of cost implications for the health system.
- In Colombia, PAVE’s HEA explore the cost implications of introducing various radical cure scenarios into the health system. A Pilot Educational Program (PEP) is evaluating the effectiveness of training health personnel at all levels in diagnosing G6PD deficiency. Both HEA and PEP results are expected for Q1 2023.
- In Peru, the Ministry of Health worked with PAVE to develop and approve a strategy for the potential introduction of TQ and quantitative G6PD testing following the implementation of a feasibility study, which plans to begin patient recruitment in Q1 2023.
- PAVE has also been working to identify challenges in case management in Guatemala, Honduras and Panama.

PAVE consolidates project work from multiple funders and is aligned with country partners and the WHO to accelerate progress. The partnership combines investments from Unitaid, the Bill & Melinda Gates Foundation, the UK Foreign, Commonwealth and Development Office (FCDO) and MMV core funding, among others. For additional information regarding our work, reach out to: pave@path.org