Laboratory Preparedness and Response Branch
Biological Response Section

Specimen Requirements for *Plasmodium* spp. Detection and Identification by Real-Time (RTi) Polymerase Chain Reaction (PCR) on clinical specimens.

Methodology: *Plasmodium* spp. (*vivax, falciparum, malariae, and ovale*) RTi-PCR

Performed: *Plasmodium* spp. RTi-PCR is performed only on specimens approved by the Disease Investigation Branch (DIB) of the Disease Outbreak and Control Division (DOCD), Department of Health (DOH).

Criteria for testing: Clinical signs and symptoms compatible with *Plasmodium* spp. infection and/or specimens meeting the case definition set by the DIB of the DOCD of the DOH.

Turn-Around-Time: Results are reported 2-3 business days after approval and receipt of specimen(s).

Specimen type required: Blood.

A minimum of 200uL sample is required, 500uL is the ideal sample quantity. Formalin fixed specimens are not suitable for molecular studies.

Specimen storage/transport: Collect a 1-5mL blood samples in Vacutainer® EDTA tubes prior to anti-parasitic therapy and ship at 4°C. Samples will be stored at the temperature it was received.

Follow packing and shipping instructions of the U.S. Department of Transportation (U.S. DOT), and the International Air Transport Association (IATA) for Biological Substance, Category B.

Specimen submission: Specimens are submitted from requesting laboratories with 81.3 submission form(s). Test is subject to CLIA regulation which require two patient identifiers on the specimen container and the test requisition(s).

The Epidemiology Specialist of the DIB must notify the Biological Response Section at 453-5993 or 453-5984 prior to the submission of specimens.

Criteria for rejection: Rejection criteria are an accreditation requirement and are intended to ensure specimen integrity prior to testing.
1. Specimen leaked in transit.
2. Specimen is not collected in a proper container or special handling instructions are not followed. Submitter will be asked to submit another specimen.

3. Specimen not received on cold packs (i.e. 4°C).

4. Specimen quantity is insufficient to perform the tests.

5. Unlabeled or incomplete specimen labeling and documentation. Submitter(s) will be notified to provide correct information and corrected State Laboratory Division (SLD) Form 81.3 in person or submit a written documentation by fax or e-mail.

6. Incomplete requisition form (e.g., travel history, medical history, previous lab results, if appropriate, etc.). Submitter must complete requisition form.

7. Specimen label does not match the requisition. Submitters must correct the discrepancy.

**Stability:**
If specimen cannot be transported to the SLD within 24 hours after collection, keep refrigerated then ship at 4°C.

**Requisition Form:**
Each specimen submitted must have a completed SLD Form 81.3. Submitter is responsible for completing SLD Form 81.3 including but not limited to the following information: at least two patient unique identifiers, specimen site/specimen type, date of onset, date of collection, date shipped/sent to the SLD, test(s) requested, name and address of submitter and other pertinent information.

Requisition forms shall be placed in a separate bag and shall not be packed with the specimen(s).

**Normal Value:**
No *Plasmodium spp.* (*vivax*, *falciparum*, *malariae*, and/or *ovale*) nucleic acid detected.

**Result Notification:**
Laboratory reports are reported to the submitters via fax. Laboratory reports to DIB will be posted to the DOCD Sharepoint and notified by email.

**Test performed at:**
Biological Response Section (BRS)
Laboratory Preparedness and Response Branch (LPRB)
State Laboratories Division
Department of Health
2725 Waimano Home Road
Pearl City, Hawaii 96782

**Contact:**
Remedios Gose at (808) 453-5993 or (808) 554-9992 or
Cheryl Daquip at (808) 453-5984 or (808) 453-6641
Reviewed By:

Remedios B. Gose, MSPH, RM (NRCM)
Laboratory Preparedness and Response Branch Chief

Approved By:

Edward Desmond, Ph.D.
Administrator, State Laboratories Division