Laboratory Preparedness and Response Program
Biological Response Section

Specimen Requirements for Real-Time (rti) Reverse-Transcriptase Polymerase Chain Reaction (RT-PCR) for Dengue Virus Serotypes 1-4 in Clinical Samples

Methodology:
CDC Dengue (DENV-1-4) Real-Time RT-PCR Assay

Performed:
The CDC DENV-1-4 Real-Time RT-PCR Assay is an FDA approved assay developed by the CDC for In-Vitro Diagnostic use (IVD).

Criteria for testing:
Clinical signs and symptoms compatible with Dengue virus infection and/or specimens meeting the case definition set by the Disease Investigation Branch (DIB) of the Disease Outbreak Control Division (DOCD) of the Department of Health (DOH).

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

Specimen type required:
Venous blood sample: Follow device manufacturer’s instructions for proper serum or plasma collection and separation. Serum is the preferred specimen. The best type of tube is serum separator (tiger/speckled-top). Red-top (no additives) is also acceptable. For plasma specimens, use sodium citrate collection tube. Do not use heparin (green top) or EDTA (purple top).

A minimum of 0.5 ml serum or plasma is required.

Dengue specimens must be collected within the first eight (8) days from onset of signs and symptoms.

Specimen storage / transport:
Refrigerate serum or plasma at 4°C or maintain on ice for no longer than 24 hours. If storage/transport will exceed 24 hours, freeze serum or plasma at -20°C or lower.

Ship separated serum or plasma on cold packs (i.e. 4°C) within 24 hours. Ship on dry ice if the specimens have been frozen at -20°C or lower.

Follow instructions for Class B - Biological Substance of the U.S. Department of Transportation (U.S. DOT) and
Specimen submission:
Submitters: Clinical laboratories and the Disease Investigation Branch (DIB).
Please notify DIB and the SLD Biological Response Section (BRS) prior to the submission of specimens.

Criteria for rejection:
- Specimen is received in a container that is leaking;
- Specimen is not collected in a proper container or special handling instruction is not followed;
- Blood collected with heparin or EDTA tubes;
- Specimen is not received at 4°C or packed in blue ice;
- Frozen specimens not shipped in dry ice;
- Specimen quantity is insufficient to perform the tests;
- Unlabeled specimens;
- Incomplete specimen labeling/documentation.

Stability:
Separated serum or plasma must be refrigerated at 2-8°C for no longer than 24 hours. If the specimen cannot be transported to the State Laboratories Division within 24 hours after collection and separation, it should be frozen at -20°C or lower. Frozen samples should be shipped in dry ice.

Requisition Form:
Each specimen submitted must have a completed Form 81.3. Submitter is responsible for completing SLD Form 81.3 (including but not limited to the following information: patient identifier, date of onset of illness, signs and symptoms, travel history, immunization history, name and address of submitter).

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

Normal Value:
No Dengue Virus Nucleic Acid Detected

Result Notification:
Laboratory results are reported to the submitters and Disease Investigation Branch (DIB) of the DOH Disease Outbreak Control Division (DOCD). Laboratory reports for the DIB will be posted to the DOCD SharePoint.

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

Contact: Rebecca Sciulli M.Sc., MT at 808-453-5993 or
808-368-3373
Remedios Gose at 808-453-5984

Approved By:

A. Christian Whelen, Ph.D.
Administrator, State Laboratories Division

8/5/2014 Date