

Dengue virus type 1 (DENV-1) TaqMan (DENV1-TM2017) assay

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Abstract

This protocol was designed and developed at this laboratory.

The assay specifically targets the 3' UTR region of DENV-1 and is designed as a qualitative screening test for human cases of DENV-1 infection, but not other known DENVs.

Citation: Ian Mackay, Judy Northill Dengue virus type 1 (DENV-1) TaqMan (DENV1-TM2017) assay. protocols.io

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Before start

- If using a different brand or model of real-time thermocycler, check the concentration of ROX is adequate.
- Method assumes the user is familar with the thermocycler and software used to run the protocol and with PCR in general..

Materials

SuperScript™ III Platinum™ One-Step qRT-PCR Kit <u>11732088</u> by <u>Life Technologies</u>

Protocol

Oligonucleotide sequences

Step 1.

| Name | Sequence (5'-3') |
|------------------|--|
| DENV1-2017_For | AGGAGACCCCTCCC N A |
| DENV1-2017_Rev | CCTCTAACCTCTAGTCCTTACCAC |
| DENV1-2017_Probe | 6FAM - CCCAACACCAGGGGAAGCTGTA Y CCT - BHQ |

Reagents

Step 2.



SuperScript™ III Platinum™ One-Step qRT-PCR Kit <u>11732088</u> by <u>Life Technologies</u>

Reaction set-up

Step 3.

MIX PREPARATION

| Reagent | Vol (µl) x1 | Final reaction concentration |
|-------------------------------------|-------------|------------------------------|
| Nuclease-free water | 4.41 | N/A |
| DENV1-2017_For (200μM) | 0.09 | 900nM |
| DENV1-2017_Rev (200μM) | 0.05 | 500nM |
| DENV1-2017_Probe (100μM) | 0.01 | 50nM |
| 2X Reaction Mix ¹ | 10 | 1X |
| ROX reference dye (25μM) | 0.04 | 50nM |
| SuperScript™ III/Platinum™ Taq Mix¹ | 0.4 | N/A |
| TOTAL | 15 | N/A |

¹Superscript[™]III Platinum[™] One-step qRT-PCR kit

- Dispense 15µL to each reaction well.
- Add 5µL of template (extracted RNA, controls or NTC [nuclease-free water]).
- Total reaction volume is 20µL

Amplification

Step 4.

| 50°C | 5min | 1X |
|------|--------------------|-----|
| 95°C | 2min | 1X |
| | | |
| 95°C | 3sec | 40X |
| 60°C | 30sec ¹ | |

¹Florescence acquisition step

Result analysis

Step 5.

The definition used for a satisfactory positive result from a real-time fluorogenic PCR should include each of the following:

- 1. A **sigmoidal curve** the trace travels horizontally, curves upward, continues in an exponential rise and followed by a curve towards a horizontal plateau phase
- 2. A **suitable level of fluorescence** intensity as measured in comparison to a positive control (y-axis)
- 3. A **defined threshold (C_T) value** which the fluorescent curve has clearly exceeded (Fig.1 arrow), which sits early in the log-linear phase and is <40 cycles

A flat or non-sigmoidal curve or a curve that crosses the threshold with a $C_T > 40$ cycles is considered a negative result. NTCs should not produce a curve

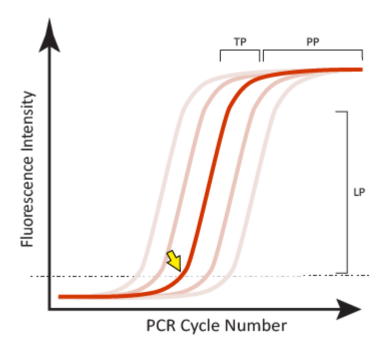


Figure 1. Examples of satisfactory sigmoidal amplification curve shape when considering an assay's fluorescent signal output. The crossing point or threshold cycle (C_T) is indicated (yellow arrow); it is the value at which fluorescence levels surpass a predefined (usually set during validation, or arbitrary) threshold level as shown in this normalized linear scale depiction. LP-log-linear phase of signal generated during the exponential part of the PCR amplification; TP-a slowing of the amplification and accompanying fluorescence signal marks the transition phase; PP-the plateau phase is reached when there is little or no increase in fluorescent signal despite continued cycling.