



# Influenza A H3 virus TaqMan assay V.3

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#### ABSTRACT

This test is a modification to the World Health Organization's influenza A H3 TaqMan documented in 'WHO information for molecular diagnosis of influenza virus - update 1' (see file below). New primers were added and the WHO primers discarded. The test targets the hemagglutinin (HA) gene.

Influenza\_WHO\_update\_2 01403rev201505.pdf

STEPS MATERIALS

NAME ✓ CATALOG # ✓ VENDOR ✓
SuperScript™ III Platinum™ One-Step qRT-PCR Kit 11732088 Life Technologies

# BEFORE STARTING

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

# Oligonucleotides

Name	5'-3' SEQUENCE
H3hFor1	GGTACGGYTTCAGGCAT
H3hRev1	TCAATCTGATGGAATTTCTCGTTG
H3h-1144dProbe	6FAM-CTGCTGCTCTCCCT-BHQ1

- The oligoprobe is from the World Health Orgnization protocol linked below.
- New primers were designed to improve assay performance.
   <a href="http://www.who.int/entity/influenza/gisrs\_laboratory/molecular\_diagnosis\_influenza\_virus\_humans\_update\_201403rev20\_1505.pdf?ua=1\_">http://www.who.int/entity/influenza/gisrs\_laboratory/molecular\_diagnosis\_influenza\_virus\_humans\_update\_201403rev20\_1505.pdf?ua=1\_</a>



### Reaction Set-up

- 3 The assay has been used on both a Rotor-Gene 6000 and a Rotor-Gene Q real-time thermocycler
  - Prepare sufficient mix for the number of reactions.
  - Include a suitable 'dead volume' as necessary if using a robotic dispenser.

Reagent	Vol (µL) X1	Final reaction concentration
Nuclease-free water	4.43	
H3hFor1 (200pmol/µL)	0.05	500nM
H3hRev1 (200pmol/µL)	0.05	500nM
H3h-1144dProbe (100pmol/μL)	0.03	150nM
2X Reaction Mix <sup>1</sup>	10.0	1X
ROX reference Dye (25µM) <sup>1,2</sup>	0.04	50nM
SuperScript <sup>TM</sup> III/Platinum <sup>TM</sup> Taq Mix <sup>1</sup>	0.4	
TOTAL	15	

- 1-Superscript<sup>TM</sup>III Platinum<sup>TM</sup> One-step qRT-PCR kit; 2-See Guidelines
- Dispense 15µL to each reaction well.
- $\,\bullet\,$  Add 5µL of template (extracted RNA, controls or NTC [nuclease-free water] ).
- Total reaction volume is 20μL

### **Amplification**

#### 4 RT-PCR

50°C	5min	
95°C	2min	
95°C	3s	40X
60°C	30s*	

<sup>\*</sup>Florescence acquisition step

#### Result analysis

- 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<sub>T</sub>) value which the fluorescent curve has clearly exceeded (Fig.1 arrow) and which sits early in the loglinear phase and is <40 cycles
  - 4. A flat or non-sigmoidal curve or a curve that crosses the threshold with a C<sub>T</sub> value >40 cycles is considered a negative result
  - 5. NTCs should not produce a curve

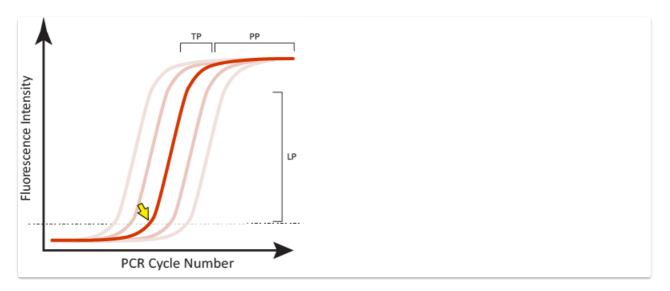


Figure 1. Examples of satisfactory sigmoidal amplification curve shape when considering an assay's fluorescent signal output. The crossing point or threshold cycle  $(C_T)$  is indicated (yellow arrow); it is the value at which fluorescence levels surpass a predefined (usually set during validation, 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.

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