

Purification of Viral RNA from Stabilized Saliva with the Maxwell[®] HT Viral TNA Kit, Custom

Purify viral RNA from stabilized saliva using the Maxwell[®] HT Viral TNA Kit, Custom with the KingFisher[™] Flex Purification System.

Kit:	Maxwell [®] HT Viral TNA Kit, Custom (Cat.# AX2340)	This protocol was developed by
Analyses:	RT-qPCR for detection of Respiratory Syncytial Virus (RSV) and Influenza B	Promega Applications Scientists and is intended for research use only. Users are responsible for determining suitability of the
Sample Type(s):	Stabilized saliva stored in Oragene∙RNA (RE-100) or Oragene∙DNA (OG-500) tubes (DNA Genotek)	protocol for their application. For further information, contact Technical Services at:
Input:	200μΙ	techserv@promega.com

Materials Required:

- Maxwell[®] HT Viral TNA Kit, Custom (Cat.# AX2340)
- 4/40 Wash Solution (Cat.# A2221)
- Alcohol Wash, Blood (Cat.# MD1411)
- 100% Ethanol and 80% Ethanol
- 100% Isopropanol
- KingFisher[™] Flex Purification System (ThermoFisher Scientific, Cat.# 24074431)
- KingFisher Deep Well 96 Plate (ThermoFisher Scientific, Cat.# 95040450)
- KingFisher 96 tip comb for DW magnets (ThermoFisher Scientific, Cat.# 97002534)
- KingFisher[™] Flex Run Protocol (Maxwell_HT_Viral_TNAv1_2_RT Elution.bdz)

Protocol:

- 1. Prepare 4/40 Wash Solution and Alcohol Wash as indicated on the bottles.
- 2. Prepare KingFisher[™] plates:
 - a. Tip Plate: Add Tip comb for KingFisher Deep Well plate.
 - b. Elution Plate: Add 110µl of Nuclease-Free Water per well.
 - c. 4_40 Wash 1: Add 900µl of 4/40 Wash Solution per well.
 - d. Alcohol Wash 2: Add 450µl of Alcohol Wash per well.
 - e. Ethanol Wash 3: Add 450µl of 80% ethanol per well.
 - f. Lysis and Bind: Add the following reagents to each well*:
 - i. 200µl of Lysis Buffer
 - ii. 35µl of Proteinase K
 - *Optional: A master mix of the above reagents (plus at least 10% overage) can be prepared immediately before sample addition and 235µl added to each well.

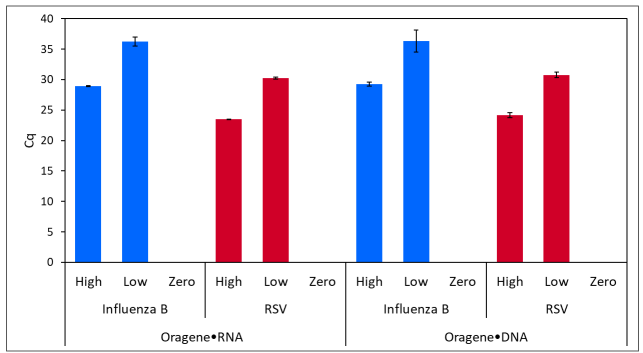


Product Application

- 3. Transfer 200µl of stabilized saliva to each well of the Lysis and Bind Plate.
- 4. Start the KingFisher[™] Flex Run Protocol (Maxwell_HT_Viral_TNAv1_2_RT Elution.bdz).
- 5. Load the KingFisher[™] 96 Deep Well Plates as directed by the instrument software.
- 6. After the heated lysis step, add 530µl of 100% Isopropanol and 35µl of MagneSil[®] RED resin (vortex vigorously to resuspend prior to addition)[^].
 ^Optional: A master mix of the above reagents (plus at least 10% overage) can be prepared and 565µl added to each well. Vortex master mix vigorously before adding to wells.
- 7. Continue the KingFisher[™] Flex Run Protocol until complete.



Product Application



Results:

Detection of RSV and Influenza B RNA extracted from stabilized saliva. Saliva was collected in Oragene•RNA (RE-100) or Oragene • DNA (OG-500) tubes (DNA Genotek) from four individuals and incubated overnight at room temperature. Stabilized saliva from each tube type was pooled. RSV A and Influenza B (Hong Kong) virus were reconstituted from Helix Elite[™] Inactivated Standard Inactivated Influenza A/B and Respiratory Syncytial Virus (Microbiologics Cat.# HE0044N) and spiked into stabilized saliva from each tube type. High virus sample contains approximately 2 x 10⁵ copies of Influenza B and RSV A per 200µl sample. Low virus sample is a 1:100 dilution of the high virus sample in stabilized saliva. 200µl of the spiked stabilized saliva was processed with the Maxwell® HT Viral TNA Kit, Custom on the KingFisher™ Flex Purification System as described above. Following nucleic acid purification, presence of RSV A and Influenza B was detected by RT-qPCR using GoTag[®] 1-Step Probe RT-qPCR System (Cat.# A6121). Each reaction contained 5µl of eluate with 12.5µl of the GoTag[®] Probe gPCR Master Mix with dUTP, 0.5µl of GoScript[™] RT Mix for 1-Step RT-qPCR, 1000nM forward and reverse primers and 200nM probe for RSV¹ or Influenza B², and Nuclease-Free Water added to a final volume of 25µl. 1-step RT-qPCR thermal cycling was as follows²: reverse transcription at 50°C for 30 minutes, hot-start activation at 95°C for 2 minutes, and then 45 cycles of denaturation at 95°C for 15 seconds and annealing/extension at 55°C for 30 seconds, with signal acquisition during the annealing/extension stage of cycling. Data represent the average of duplicate purifications amplified in duplicate. Error bars indicate the standard deviation.

References:

- 1. Fry, A.M., *et al.*, (2010) The Burden of Hospitalized Lower Respiratory Tract Infection due to Respiratory Syncytial Virus in Rural Thailand, *PLoS One*. *5*, e15098.
- 2. Selvaraju, S.B., *et al.*, (2010) Evaluation of Three Influenza A and B Real-Time Reverse Transcription-PCR Assays and a New 2009 H1N1 Assay for Detection of Influenza Viruses, *Journal of Clinical Microbiology.* **48**, 3870-3875.