

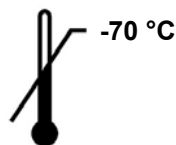
# *pro*PARAFLU<sup>TM</sup> plus<sup>+</sup>

## Instructions for Use

***For detection and discrimination of Parainfluenza 1 Virus,  
Parainfluenza 2 Virus and Parainfluenza 3 Virus***



100



**REF**  
H81VK00

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Intended Use

The ProParaflu™+ Assay is a multiplex Real-Time PCR (RT-PCR) *in vitro* diagnostic test for the qualitative detection and discrimination of Parainfluenza 1 Virus, Parainfluenza 2 Virus and Parainfluenza 3 Virus (HPIV-1, HPIV-2 and HPIV-3) nucleic acids isolated and purified from nasopharyngeal (NP) swab specimens obtained from individuals exhibiting signs and symptoms of respiratory tract infections. This assay targets the conserved regions of the Hemagglutinin-Neuraminidase (HN) gene of HPIV-1, HPIV-2 and HPIV-3, respectively. The detection and discrimination of HPIV-1, HPIV-2 and HPIV-3 nucleic acids from symptomatic patients aid in the diagnosis of human respiratory tract parainfluenza infections if used in conjunction with other clinical and laboratory findings. This test is not intended to detect Parainfluenza 4a or Parainfluenza 4b Viruses.

Negative test results are presumptive and should be confirmed by cell culture. Negative results do not preclude Parainfluenza 1, 2 or 3 virus infections and should not be used as the sole basis for treatment or other management decisions.

 Summary and Explanation

Human Parainfluenza viruses are negative-sense single stranded RNA viruses surrounded by fusion protein and hemagglutinin–neuraminidase glycoprotein “spikes” on the surface<sup>1</sup>. There are four serotypes of HPIV (1 through 4). HPIVs are the second major causative agents of lower respiratory tract infections in infants and young children<sup>2</sup>. Symptoms of infection with HPIV include common cold with fever, croup, bronchiolitis, and pneumonia<sup>3</sup>. Re-infections with HPIVs are common throughout life, especially in elderly and immunocompromised patients. Each of the four serotypes has distinct clinical and epidemiological features. HPIV-1 and HPIV-2 are the leading causes of croup in children while HPIV-3 is more often associated with bronchiolitis and pneumonia. HPIV-4 has a low recovery rate in cell culture, reportedly causes mild respiratory disease and historically has not been included in the routine respiratory virus testing in most clinical virology laboratories<sup>4</sup>. The incubation period for HPIVs is usually 1 to 7 days<sup>3</sup>. Transmission of HPIVs occurs through spread of respiratory secretions from infected persons or contact with contaminated surfaces or objects. HPIVs can remain infectious in aerosols for at least an hour. Internationally, HPIV has a worldwide distribution and epidemics are known to occur, particularly with HPIV-1. Approximately 41,000 individuals per year are admitted to the hospital in the U.S. for parainfluenza infections<sup>5</sup>.

## Principles of the Procedure

The ProParaflu+ Assay enables detection and differentiation of Parainfluenza 1 Virus, Parainfluenza 2 Virus, Parainfluenza 3 Virus and an Internal Control.

An overview of the procedure is as follows:

1. Collect nasopharyngeal swab specimens from symptomatic patients using a polyester, rayon or nylon tipped swab and place into viral transport medium (refer to **Materials Required but not Provided section of this Instructions for Use.**).
2. Add an Internal Control (IC) to every sample to monitor for inhibitors present in the specimens.
3. Perform isolation and purification of nucleic acids using a MagNA Pure LC System (Roche) and the MagNA Pure Total Nucleic Acid Isolation Kit (Roche) or a NucliSENS easyMAG System (bioMérieux) and the Automated Magnetic Extraction Reagents (bioMérieux).
4. Add purified nucleic acids to ProParaflu+ Supermix along with enzymes included in the ProParaflu+ Assay Kit. The ProParaflu+ Supermix contains oligonucleotide primers and target-specific oligonucleotide probes. The primers are complementary to highly conserved regions of genetic sequences for these respiratory viruses. The probes are dual-labeled with a reporter dye attached to the 5'-end and a quencher dye attached to the 3'-end (see table below).
5. Perform reverse transcription of RNA into complementary DNA (cDNA) and subsequent amplification of DNA in a Cepheid SmartCycler II instrument. In this process, the probe anneals specifically to the template followed by primer extension and amplification. The ProParaflu+ Assay is based on Taqman reagent chemistry, which utilizes the 5' – 3' exonuclease activity of the Taq polymerase to cleave the probe thus separating the reporter dye from the quencher. This generates an increase in fluorescent signal upon excitation from a light source. With each cycle, additional reporter dye molecules are cleaved from their respective probes, further increasing fluorescent signal. The amount of fluorescence at any given cycle is dependent on the amount of amplification product present at that time. Fluorescent intensity is monitored during each PCR cycle by the real-time instrument.

Analyte	Gene Targeted	Probe Fluorophore	Absorbance Peak	Emission Peak	Instrument Channel
Parainfluenza 1 Virus	Hemagglutinin neuraminidase	FAM	495 nm	520 nm	FAM
Parainfluenza 3 Virus	Hemagglutinin neuraminidase	CAL Fluor Orange 560	540 nm	561 nm	TET
Parainfluenza 2 Virus	Hemagglutinin neuraminidase	CAL Fluor Red 610	595 nm	615 nm	Texas Red
Internal Control	NA	Quasar 670	647 nm	667 nm	Cy5

**Materials Provided**
**ProParaflu+ Assay Kit (Cat. # H81VK00)**

Reagents	Description	Quantity/ Tube	Cap Color	Cat. #	Reactions/ Tube
ProParaflu+ Supermix	<ul style="list-style-type: none"> <li>➤ Taq DNA polymerase</li> <li>➤ 4 oligonucleotide primer pairs</li> <li>➤ 4 oligonucleotide probes</li> <li>➤ Buffer containing dNTPs (dATP, dCTP, dGTP, dTTP),</li> <li>➤ MgCl<sub>2</sub> and stabilizers</li> </ul>	1030 µL	Brown	HSM81	50  (2 tubes provided)
M-MLV Reverse Transcriptase II	<ul style="list-style-type: none"> <li>➤ 11.4 U/µL</li> </ul>	36 µL	White	GLS32	100
RNase Inhibitor II	<ul style="list-style-type: none"> <li>➤ 40 U/µL</li> </ul>	120 µL	Green	GLS33	100
Positive Control (PC) - Parainfluenza RNA Control	<ul style="list-style-type: none"> <li>➤ Non-infectious <i>in vitro</i> transcribed RNA of specific viral sequences</li> </ul>	500 µL	Red	HCT81	25
Internal Control (IC) - Internal RNA Control III	<ul style="list-style-type: none"> <li>➤ Non-infectious <i>in vitro</i> transcribed RNA</li> </ul>	30 µL	Yellow	GCT12	100

**Materials Required But Not Provided**
*Plasticware and consumables*

- Polyester, rayon or nylon tipped nasopharyngeal swabs
- RNase/DNase-Free 1.5 mL polypropylene microcentrifuge tubes
- Sterile RNase/DNase-free filter or positive displacement micropipettor tips
- MagNA Pure LC System Disposables (Reagent Tubs, Reaction Tips, Tip Trays, Cartridges) or easyMAG Disposables (Sample Vessels and Tips)
- Biohit Pipette Tips for use with easyMAG System
- Greiner Break Four uncoated plate for use with easyMAG System
- Cepheid PCR reaction tubes, 25 µL
- Parafilm M or MagNA Pure LC Cartridge Seals

*Reagents*

- Roche MagNA Pure LC Total Nucleic Acid Isolation Kit (*Roche Cat. No.03038505001*) for 192 isolations or bioMérieux NucliSENS easyMAG reagents (*Buffer 1 Cat. No. 280130, Buffer 2 Cat. No. 280131, Buffer 3 Cat. No. 280132, Magnetic Silica Cat. No. 280133, and Lysis Buffer Cat. No. 280134*)
- Micro Test M4 Viral Transport Medium (*Remel, Inc. Cat. No. R12500*), Micro Test M5 Viral Transport Medium (*Remel, Inc. Cat. No. R12515*), Micro Test M6 Viral Transport Medium (*Remel, Inc. Cat. No. R12530*), Micro Test M4RT Viral Transport Medium (*Remel, Inc. Cat. No. R12505*), Copan Universal Transport Medium (Copan Diagnostics, Inc., *Cat. No. 330C*), or BD Universal Viral Transport vial, 3mL (*Becton, Dickinson and Co. Cat. No. 220220*)
- Molecular Grade Water (*RNase/DNase Free*)

### Equipment

- – 70°C Freezer
- Roche MagNA Pure LC System with Software version 3.0.11 or bioMérieux NucliSENS easyMAG System with Software version 1.0.1 or 2.0
- Biohit multi-channel pipettor for use with easyMAG
- bioMérieux NucliSENS easyMAG Instrument
- Cepheid SmartCycler II Real Time Instrument with Dx Software version 1.7b, 3.0a or 3.0b
- Micropipettes (range between 1-10 µL, 10-200 µL and 100-1000 µL)
- Mini-centrifuge with adapter for Cepheid Reaction Tubes
- Cepheid cooling block

## Warnings and Precautions

- For *in vitro* diagnostic use only.
- Performance characteristics of this assay have only been determined with nasopharyngeal swab specimens.
- Use of this product should be limited to personnel who have been trained in the techniques of Real-Time PCR.
- Handle all specimens as if infectious using safe laboratory procedures such as those outlined in CDC/NIH *Biosafety in Microbiological and Biomedical Laboratories* and in the CLSI Document M29 *Protection of Laboratory Workers from Occupationally Acquired Infections*. Thoroughly clean and disinfect all surfaces with 10% bleach. Autoclave any equipment or materials that have contacted clinical specimens before discarding.
- Use micropipettes with aerosol barrier or positive displacement tips for all procedures.
- Always pre-plan, organize and segregate workflow. Workflow in the laboratory should proceed in a unidirectional manner, beginning in the Pre-Amplification Area and moving to the Amplification/Detection Area.
  - Begin pre-amplification activities with reagent preparation and proceed to specimen preparation.
  - Always dedicate supplies and equipment to a specified area; no cross-movement allowed between areas.
  - Do not use equipment and supplies used for reagent preparation for specimen preparation activities or for pipeting or processing other sources of target nucleic acid.
  - Keep all amplification supplies and equipment in the Amplification/Detection Area at all times.
  - Always wear disposable gloves in each area and change them before entering a different area.
  - Do not open sample tubes following PCR.
- Take care to preserve the purity of kit reagents. Avoid contamination from Positive Controls and specimens by following good laboratory practices.
- Do not use kit after its expiration date.
- Do not mix reagents with different lot numbers or substitute reagents from other manufacturers.
- Material Safety Data Sheets (MSDS) are available on the Gen-Probe Prodesse, Inc. website at [www.prodesse.com](http://www.prodesse.com) from the **Technical Information** tab.

## Reagent Storage, Handling and Stability

- ⇒ Store all reagents (opened and unopened) at  $\leq -70^{\circ}\text{C}$ .
- ⇒ Always check the expiration date on the reagent tubes. For Intermediate stock of the Internal Control, use the expiration date of the originating stock control tube. Do not expose controls to more than one (1) freeze-thaw cycle.
- ⇒ ProParaflu+ components are shipped frozen, should arrive frozen and should be stored frozen after receipt. If the kit contents are not frozen, contact Gen-Probe Prodesse, Inc. for assistance.
- ⇒ An internal study demonstrated that performance of ProParaflu+ Supermix, M-MLV Reverse Transcriptase II, and RNase Inhibitor II are not affected for up to 5 freeze-thaw cycles.
- ⇒ Visually examine reagents for adequate reagent volume before beginning any test procedures.
- ⇒ Protect the ProParaflu+ Supermix from light.
- ⇒ **Controls and aliquots of controls must be thawed and kept on ice during preparation and use.**

## Specimen Collection, Handling and Storage

### Collecting the Specimen

To obtain nasopharyngeal swab samples:

1. Insert a flexible-shaft polyester, rayon or nylon tipped swab containing a dry tip into one nostril and into the nasopharyngeal area.
2. Press the swab gently against the nasopharyngeal wall to allow the swab to absorb secretions.
3. Rotate the swab two to three times and withdraw it.
4. Place the swab into a tube containing 3 mL of viral transport medium (Remel M4, M4RT, M5, M6; Copan UTM; or Becton Dickenson UVT).
5. Break off the shaft of the swab and cap the tube.



**Note**

Using a smaller volume of the viral transport medium may result in inhibition.

### Transporting Specimens

Ensure that when transporting human respiratory specimens, all applicable regulations for the transport of etiologic agents are met. Transport human respiratory specimens refrigerated at  $2-8^{\circ}\text{C}$ .

### Storing Specimens

Store specimens refrigerated ( $2-8^{\circ}\text{C}$ ) for up to 72 hours before processing. Store any leftover specimens at  $\leq -70^{\circ}\text{C}$ . If retesting a frozen specimen, thaw specimen quickly (1 to 2 minutes) in a  $37^{\circ}\text{C}$  water bath and immediately place on ice or thaw specimen on ice.

### Storing Purified Nucleic Acid

Store purified nucleic acids at  $\leq -70^{\circ}\text{C}$ . They should be tested after no more than one (1) freeze-thaw cycle.



**Note**

*Inadequate or inappropriate specimen collection, storage and transport are likely to yield false negative results.*



**Recommendation**

*Training in specimen collection is highly recommended because of the importance of specimen quality.*

## Reagent and Control Preparation

### Reagents



Note

Prepare reagents from the Roche MagNA Pure LC Total Nucleic Acid Isolation Kit or the bioMérieux Automated Magnetic Extraction Reagents following the manufacturer's instructions.

### Controls



Recommendation

- ❖ For aliquots of the Positive Control and Intermediate Stock of the Internal Control, use the expiration date of the originating stock control tube.
- ❖ Controls and aliquots of controls must be thawed and kept on ice at all times during preparation and use.

### Positive Control (PC)



Note

Include the Positive Control (the red cap tube) with each RT-PCR run.

1. Thaw Positive Control on ice.
2. Make 25 aliquots of 20  $\mu\text{L}$ , label and store at  $\leq -70^{\circ}\text{C}$ . Ensure that aliquots do not undergo more than one (1) freeze-thaw cycle.
3. Dilute the Positive Control just prior to setup of the RT-PCR reaction (see **Step 4 (a)** of the **Assay Procedure**).



Recommendation

Do not spike Positive Control with the Internal Control. Do not take Positive Control through the nucleic acid isolation procedure.

### Internal Control (IC)

1. Thaw Internal Control (the yellow cap tube) on ice.
2. Create Intermediate stock tubes of the Internal Control using the following dilution scheme:

$$26 \mu\text{L Internal Control} + 65 \mu\text{L RNase Inhibitor} + 2509 \mu\text{L molecular grade water} = 2600 \mu\text{L total volume}$$

3. Make aliquots of 110  $\mu\text{L}$ , label, and store at  $\leq -70^{\circ}\text{C}$  (this is enough volume to add to 5 samples at 20  $\mu\text{L}$  per sample). Make aliquots of larger or smaller volumes based on the number of samples expected to be processed in a single run. Ensure that aliquots do not undergo more than one (1) freeze-thaw cycle.
4. Add the appropriate volume of Intermediate stock of the Internal Control to each sample prior to nucleic acid isolation (see **Step 1** of the **Assay Procedure**).
5. **Save remaining RNase Inhibitor by re-freezing the leftover volume in the original tube for use in Step 4 of the Assay Procedure.**

### Negative Control (NC)

1. Use Viral Transport Medium as the Negative Control.
2. Add the appropriate volume of Intermediate stock of the Internal Control to the Negative Control prior to nucleic acid isolation (see **Step 1** of the **Assay Procedure**).

### Extraction Control (EC)

Good laboratory practice recommends including a positive extraction control (e.g. previously characterized positive sample or negative sample spiked with a well characterized HPIV-1, HPIV-2 or HPIV-3 strain) in each nucleic acid isolation run. The extraction control should be treated as a sample during assay performance and analysis.

**Assay Procedure**
**Assay Overview:**

**Get Ready:** Create the Assay Protocol for the Cepheid SmartCycler instrument using the Dx Software (first time only).

1. **Prepare the Samples and Negative Control.**
2. **Isolate the Nucleic Acid** – MagNA Pure LC System using the Total Nucleic Acid Isolation (TNAI) Kit  
**OR**
3. **Isolate the Nucleic Acid** – NucliSENS easyMAG System using the Automated Magnetic Extraction Reagents.
4. **Set up the RT-PCR Reaction.**
5. **Run the ProParaflu+ Assay.**
6. **Print report.**


**Note**

- ❖ Instructions provided for the Cepheid SmartCycler Real Time Instrument with Dx Software version 1.7b and 3.0a / 3.0b.
- ❖ **Do NOT deviate from the protocol settings defined in this section.**

**Get Ready:** Create the Assay Protocol for the Cepheid SmartCycler instrument using the Dx Software (first time only)


**Note**

- ❖ The protocol is only created for first-time use; it does not need to be recreated with each sample run.
- ❖ Refer to SmartCycler Dx Software Operator Manual for assistance in defining assay protocols.
- ❖ To **Define** and **Edit** Assay protocols, the user must have administrative access rights, otherwise the fields will be grayed out.
- ❖ Cepheid Dx Software interprets the data and reports the run as either **VALID** or **INVALID**, based on the results of the Negative Control. Enter the Positive Control and Extraction Control as if they were samples.
- ❖ Interpret the control results and determine if the run is **VALID** or **INVALID**. All Control criteria must be met in order for the run to be **VALID** (see **Interpretation of Results** section).

**1. Create the ProParaflu+ protocol:**

- a. Launch the Cepheid Dx software application.
- b. Click on the **Define Assay** box at the top of the screen.
- c. Click on the **New Assay** box at the bottom of the screen.
- d. Enter **ProParaflu+ Assay** for the assay protocol in the window that opens.
- e. Click **OK**.
- f. In the middle of the **Define Assay** screen, select **FTTC25** for the **Dye Set**.
- g. Select **NC fails if: Any target criterion is positive** (Dx 3.0a/3.0b: this is default setting in Control Settings Tab).
- h. Check the box to **Require Lot Number** (Dx 3.0a/3.0b: click on Advanced Tab and check Require Lot Number box).
- i. Deselect the box to **Use Patient IDs** (Dx 3.0a/3.0b: click on Advanced Tab, Use Patient IDs box is deselected by default).

- j. Enter **Thermocycler Parameters** in the Protocol section (bottom half of **Define Assay** screen).

Stage 1			Stage 2			Stage 3 Repeat 5 times			Stage 4 Repeat 45 times		
Hold			Hold			2- Temperature Cycle			2- Temperature Cycle		
Temp	Secs	Optics	Temp	Secs	Optics	Temp	Secs	Optics	Temp	Secs	Optics
42	1800	OFF	95	600	OFF	95	30	OFF	95	10	OFF
						55	60	ON	55	60	ON

Stages 5 – 10 remain UNUSED

2. Enter information in **Analysis Settings** tab as follows:



Gray boxes are default settings.

Note

Channel	Dye Name	Target	Usage	Curve Analysis	Thresh Setting	Manual Thresh	Auto Thresh	Auto Min. Cycle	Auto Max. Cycle	Valid Min. Cycle	Valid Max. Cycle	Bkgnd Sub	Bkgnd Min. Cycle	Bknd Max. Cycle	Boxcar Avg	EndPt Thresh	NC IC %	IC Delta
1	FAM*	HPIV-1	Assay**	Primary Curve	Manual Thresh	60	NA	5	10	13.0	50	On	5	50	0	60	10†	NA
2	TET*	HPIV-3	Assay**	Primary Curve	Manual Thresh	40	NA	5	10	13.0	50	On	5	50	0	40	10†	NA
3	TxR*	HPIV-2	Assay**	Primary Curve	Manual Thresh	50	NA	5	10	13.0	50	On	5	50	0	50	10†	NA
4	Cy5*	Internal Control	Internal Control	Primary Curve	Manual Thresh	40	NA	5	10	13.0	50	On	5	50	0	40	10†	NA

\*If the Dye Names are incorrect, check that FTTC25 Dye Set is being used.

\*\*Dx 3.0a/3.0b = Target

† Dx 3.0a/3.0b = NA

3. Enter information in the **Control Settings** tab.



Note

- ❖ Enter Positive Control and/or the Extraction Control as a sample. Do not use the Positive Control Settings. Enter 0 Replicates to inactivate Positive Controls PC1-3. A 3079 error (Fluorescence Signal Too High) in the Positive Control invalidates the run, this is avoided if the Positive Control and/or the Extraction Control is entered as a sample and the results for each of the target channels are individually evaluated.
- ❖ Use only one Negative Control (NC1). Enter 0 Replicates to inactivate the Negative Controls NC2 and NC3.
- ❖ Gray boxes are default settings.

Control ID	Control Name	Replicate	HPIV-1 Valid Min Cycle	HPIV-1 Valid Max Cycle	HPIV-1 EndPt Thresh	HPIV-3 Valid Min Cycle	HPIV-3 Valid Max Cycle	HPIV-3 EndPt Thresh	HPIV-2 Valid Min Cycle	HPIV-2 Valid Max Cycle	HPIV-2 EndPt Thresh	IC +/-	IC Valid Min Cycle	IC Valid Max Cycle	IC EndPt Thresh
PC1	Pos Cntrl 1	0	13.0	45.0	10	13.0	45.0	10	13.0	45.0	10	+	13.0	45.0	10
PC2	Pos Cntrl 2	0	13.0	45.0	10	13.0	45.0	10	13.0	45.0	10	+	13.0	45.0	10
PC3	Pos Cntrl 3	0	13.0	45.0	10	13.0	45.0	10	13.0	45.0	10	+	13.0	45.0	10
NC1	Neg Cntrl	1	13.0	50.0	60	13.0	50.0	40	13.0	50.0	50	+	15.0	50.0	40
NC2	Neg Cntrl 2	0	13.0	45.0	10	13.0	45.0	10	13.0	45.0	10	+	13.0	45.0	10
NC3	Neg Cntrl 3	0	13.0	45.0	10	13.0	45.0	10	13.0	45.0	10	+	13.0	45.0	10

#### 4. Confirm **Probe Check Settings** tab.



Note

*The probe check is not used for the ProParaflu+ Assay Protocol.*

Probe Check Settings													
Ch #	Dye Name	Prb 1 Min	Prb 1 Max	Prb 2 Min	Prb 2 Max	Prb 3 Min	Prb 3 Max	Delta 12 Min	Delta 12 Max	Delta 23 Min	Delta 23 Max	Delta 13 Min	Delta 13 Max
1	FAM	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
2	TET	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
3	TxR	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
4	Cy5	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

#### 5. Select **Save Assay**.

##### 1. Prepare the Samples, Extraction Control, and Negative Control (Pre-Amplification Area)

###### a. Add Internal Control to all samples.

- i. Thaw the appropriate number of aliquots of Intermediate stock of the Internal Control (enough volume needed for each sample, the Extraction Control, and the Negative Control) on ice.
- ii. Remove 180  $\mu$ L of sample from the original sample tube and pipet into a labeled 1.5 mL microcentrifuge tube. Alternately, pipet 180  $\mu$ L of sample directly into sample cartridge or sample vessel.
- iii. Remove 180  $\mu$ L of Extraction Control from the original sample tube and pipet into a labeled 1.5 mL microcentrifuge tube. Alternately, pipet 180  $\mu$ L of Extraction Control directly into sample cartridge or sample vessel.
- iv. Add 20  $\mu$ L of Intermediate stock of the Internal Control to each sample. **Pipet up and down a minimum of 5 times to mix** using a new pipet tip for each sample.
- v. Keep tubes on ice.
- vi. Store any remaining sample at  $\leq -70^{\circ}\text{C}$ .

###### b. Add Internal Control to the Negative Control.

- i. Include one (1) Negative Control in each run.
- ii. Add 180  $\mu$ L of Viral Transport Medium to a labeled 1.5 mL microcentrifuge tube. Alternately, pipet 180  $\mu$ L of Viral Transport Medium directly into sample cartridge or sample vessel.
- iii. Add 20  $\mu$ L of Intermediate Stock of Internal Control to the Viral Transport Medium. **Pipet up and down a minimum of 5 times to mix.**
- iv. Keep tube on ice.
- v. Discard remaining volume of Internal Control – DO NOT reuse.



*Do not reuse Internal Control.*

##### 2. Isolate the Nucleic Acid (Pre-Amplification Area) - MagNA Pure LC System using the Total Nucleic Acid Isolation (TNAI) Kit

###### a. Start the instrument and software.

- i. Turn power on to the MagNA Pure LC instrument and then turn on the computer.
- ii. Start the MagNA Pure LC software.
- iii. From the *Main Menu* screen, select **Sample Ordering** and enter sample information in **Sample Name** column.
- iv. Select the **Total NA Variable\_elution\_volume.blk** protocol.
- v. Follow the software instructions and specify the number of samples.
- vi. Type in **200  $\mu$ L** for the sample volume and verify that 50  $\mu$ L elution volume is selected.

vii. Select **Stage Setup** and the software will automatically calculate the amount of each reagent that is required.

**b. Fill the reagent tubs**

Before starting the isolation procedure, fill all reagent tubs outside the instrument with the required volume of each reagent listed on the **Start Information** screen.



Note

*Use only the reagent amount needed for the number of samples entered into the software. Reagents are not stable for long-term storage in tubs. Vortex Magnetic Glass Particles (MGPs) and load the exact amount of MGPs (as listed on the **Start Information Screen**) into the instrument just before the run starts.*

**c. Load reagent tubs and disposables into the instrument**

Use the information on the **Start Information** screen to place all disposable plastics and reagent tubs necessary for the batch run on the Reagent/Sample Stage.



Recommendation

*Use a colored "Positioning Frame" (provided with the TNAI kit) on the Reagent Tub Rack to help to correctly load reagents.*

**d. Load the Samples, Extraction Control, and Negative Control into the MagNA Pure Sample Cartridge.**

- i. Transfer 200  $\mu$ L of each sample to individual wells in the MagNA Pure Sample Cartridge.
- ii. Transfer 200  $\mu$ L of the Extraction Control and Negative Control to different wells in the MagNA Pure Sample Cartridge.
- iii. Cover cartridge with Parafilm or MagNA Pure LC Cartridge Seal and keep cartridge on ice until ready to load the instrument.

**e. Load the samples.**

Transfer cartridge containing the samples, Extraction Control, and Negative Control into the MagNA Pure LC instrument.

**f. Start the run**

- i. Start the Batch Run by confirming the correct placement of all disposable plastics and reagents by mouse-clicking the respective text boxes on the **Start Information** screen.
- ii. Select the **OK** button to start the automated isolation procedure. The instrument will automatically dispense all reagents and process the samples.

**g. Store the eluted total nucleic acid**

After completing the run, place the Storage Cartridge containing the eluted nucleic acids immediately on ice or transfer eluted nucleic acid to 1.5 mL tubes and store for longer durations at  $\leq -70^{\circ}\text{C}$ .



Note

*Do not store purified nucleic acids in the Storage Cartridge on the Cooling Unit 1.*

### 3. Isolate the Nucleic Acid (Pre-Amplification Area) - NucliSENS easyMAG System using the Automated Magnetic Extraction Reagents

#### a. Start instrument and software.

Turn power on to the easyMAG instrument and once the LED on the instrument turns green, turn on the computer and log into the software.

#### b. Prepare the software for a run.

To prepare for a run, touch the “**Settings**” icon in the main toolbar, which defaults to the “Application Settings” icon, and choose the following run settings:

**Default Request:** Generic 1.0.6 or 2.0.1 (for software version 1.0.1 or 2.0, respectively)

**Run Name Prefix:** N/A (leave as default)

**Sample ID prefix:** N/A (leave as default)

**Sample Type:** Primary (on-board lysis)

**Default On-board Lysis Dispensing:** Yes

**Default On-board Lysis Incubation:** Yes

**Sample Addition Guidance:** Off

**Reagent Tracking:** Off

#### c. Input buffer information.

Touch the “**Instrument**” icon to default to the “**Reagent Inventory**” icon and input the buffer barcodes by first scanning the instrument position (A, B, C, or D) and then its corresponding buffer. For example, scan position A and then scan the bottle of Lysis buffer in that position and then move on to position B and its corresponding bottle.

#### d. Create a worklist.

i. Touch the “**Daily Use**” icon which will default to the “**Define Extraction Request**” icon and select the following settings:

**Sample ID:** Manually enter the sample name.

**Matrix:** Other

**Request:** Generic 1.0.6 or 2.0.1 (for software version 1.0.1 or 2.0, respectively)

**Volume (mL):** 0.20 (input volume of sample)

**Elate (µL):** 55

**Type:** Primary

**Priority:** Normal

ii. Press **Enter** on the keyboard or touch the “**New Extraction Request**” icon after each manual sample addition. The settings above will remain as the default settings for each subsequent entry as long as you do not navigate to other pages.

#### e. Create a run and add samples from the worklist.

Touch the “**Organize Runs**” icon and then the “**Create Run**” icon, which will bring up the *New Run Window*. In this screen, name the run appropriately and verify that the **auto-number** box is left unchecked (NOT SELECTED) and that **Yes** is selected for both the On-Board Lysis Dispensing and On-Board Lysis Incubation options. Touch **OK** and the *New Run Window* will close and you will be in the “**Organize Runs**” screen. Assign samples to the run with the positioning (arrow) icons. Touch the “**Load Run**” icon and select the run. Print the worklist with the “**Print worklist**” icon. This worklist will aid in keeping track of the order of the samples to be loaded into the sample vessel wells.

#### f. Load samples, Extraction Control, Negative Control, and tips and barcode the samples vessel(s).

Add all of the 200 µL from each sample, Extraction Control, and Negative Control into the proper wells of the sample vessel(s) as noted in the worklist. Insert tips and sample vessel(s) in the correct order as noted in the worklist and scan the sample vessel(s) position on the instrument and then the sample vessel itself. For example, scan position A and then the sample vessel in that position, then B and then C, if necessary. After scanning the sample vessel(s), the indicator will change from red to green on the

screen.

**g. On-Board Lysis Dispensing.**

Once the samples and tips are loaded and the vessel(s) scanned, close the lid and touch the “**Dispense Lysis**” icon. The instrument will dispense 2 mL of Lysis buffer and incubate for 10 minutes.

**h. Prepare the magnetic silica to add to the sample vessel wells.**

During the 10-minute lysis incubation, use the Biohit multi-channel pipettor to prepare the magnetic silica. This procedure will need to be performed for each sample vessel used in the run (1, 2, or 3 times).

- i.** Set the pipettor to **Program 1** and place a Biohit pipette tip on position 1. Program 1 provides the means to aspirate and dispense 550  $\mu$ L of liquid. The magnetic silica is prepared in a 1:1 ratio of Molecular Biology Grade Water to Magnetic Silica.
- ii.** Using Program 1 of the pipettor, press the **start** button to aspirate and then again to dispense 550  $\mu$ L of water into a microcentrifuge tube. Vortex the tube of magnetic silica briefly to mix and use Program 1 of the pipettor to aspirate and then dispense 550  $\mu$ L of magnetic silica into the same microcentrifuge tube as the water. Eject the tip, cap the tube, and vortex to mix.
- iii.** Set the pipettor to **Program 2** and place a Biohit pipette tip on position 1. Program 2 will transfer 8 volumes of the previous mix to the 8 wells of a strip on an ELISA plate (1 strip/sample vessel). Press the **start** button to aspirate the mix. Press the **start** button again to dispense the remaining mixture back into the tube containing the mix to reset the pipette.
- iv.** Press the **start** button 8 separate times to dispense the remaining mix in each of 8 wells of an ELISA plate strip and eject the tip.
- v.** After the 10 minute lysis incubation is done, set the pipettor to **Program 3** and place 8 Biohit pipette tips on the multi-channel pipettor (or however many samples are present in the specific sample vessel). Make sure that the filter tips are very well connected with the multi-channel pipettor to prevent leakage errors. Program 3 first mixes the magnetic silica mixture in the ELISA plate and then aspirates it to be delivered to the wells of the sample vessel where it will be mixed. Press the **start** button once and the pipette will mix the silica in the ELISA plate and then aspirate it for addition to the sample vessel. Verify that each tip has the same volume of silica mix before placing in the sample vessel. Place the pipettor over the sample vessel strip so the tips are below the liquid level of each sample and press the **start** button again, which will then aspirate 800  $\mu$ L out of the sample vessel strip well and perform 3 mix cycles with 1000  $\mu$ L. At this point be sure to hold the pipette steady below the liquid/air interface as it is mixing so as not to introduce bubbles to the sample.
- vi.** Repeat for each sample vessel in the run.

**i. Start the run.**

Touch the “**Start**” icon to begin the run. The instrument will perform 5 washes and heat and elute. The purified nucleic acids need to be transferred to appropriate storage tubes (1.5 mL microcentrifuge tubes) on ice within 30 minutes of extraction completion to avoid contamination by the magnetic silica stuck to the front wall of the sample vessel(s). Use immediately or store at  $-70^{\circ}\text{C}$ .

#### 4. Set up the RT-PCR Reaction (Pre-Amplification Area)



Start the SmartCycler ProParaflu+ Assay run within 1 hour of making the RT-PCR Master mix.

Note

##### a. Dilute the Positive Control.

- i. Include the Positive Control with each run. Thaw one (1) aliquot of the Positive Control on ice.
- ii. Add 45  $\mu\text{L}$  of molecular grade water to an individual 1.5 mL microcentrifuge tube.
- iii. Transfer 5  $\mu\text{L}$  of the Positive Control to the tube. **Pipet up and down a minimum of 5 times to mix.**
- iv. Keep tube on ice.
- v. Discard remaining volume of Positive Control – DO NOT reuse.



*Do not reuse Positive Controls.*

##### b. Prepare the RT-PCR Master mix



The RT-PCR Master mix must be prepared FRESH for each RT-PCR run.

Note

- i. Calculate the amount of each reagent needed based on the number of reactions (samples + controls):

19.45 $\mu\text{L}$	ProParaflu+ Supermix
+0.30 $\mu\text{L}$	M-MLV Reverse Transcriptase
+ <u>0.25 <math>\mu\text{L}</math></u>	RNase Inhibitor
20.00 $\mu\text{L}$	per reaction

- ii. Thaw the ProParaflu+ Supermix on ice and **mix by pipetting up and down a minimum of 5 times before use.**
  - iii. Remove M-MLV Reverse Transcriptase and RNase Inhibitor enzymes from the freezer and keep on ice during use.
  - iv. Prepare the RT-PCR Master mix by combining the reagents listed above in a 1.5 mL microcentrifuge tube. **Pipet up and down a minimum of 5 times to mix.**
  - v. Keep the RT-PCR Master mix on ice and protected from light before adding to SmartCycler tubes.
- c. Add 20  $\mu\text{L}$  of RT-PCR Master mix to the SmartCycler tubes.**
- i. Load the required number of tubes into the Cepheid Cooling Block.
  - ii. Pipet 20  $\mu\text{L}$  of the RT-PCR Master mix into the upper part of the SmartCycler tubes. Discard any unused RT-PCR Master mix.
- d. Add 5  $\mu\text{L}$  of each sample's nucleic acid to individual SmartCycler tubes containing RT-PCR Master mix.**
- i. After adding the sample's nucleic acid to the SmartCycler tube, **pipet up and down 2 to 3 times in the upper part of the tube.**
  - ii. Close the tube. Use a new pipette tip for each sample.

- e. **Add 5  $\mu$ L of the diluted Positive Control to a separate SmartCycler tube containing RT-PCR Master mix.**
  - i. After adding the diluted Positive Control to the SmartCycler tube, **pipet up and down 2 to 3 times in the upper part of the tube using a new pipet tip for each control tube.**
  - ii. Close the tube.
  - iii. Discard remaining volume of diluted Positive Control – DO NOT reuse.



*Do not reuse diluted Positive Controls.*

- f. **Add 5  $\mu$ L of the Extraction Control nucleic acid to a separate SmartCycler tube containing RT-PCR Master mix.**
  - i. After adding the Extraction Control nucleic acid to the SmartCycler tube, **pipet up and down 2 to 3 times in the upper part of the tube.**
  - ii. Close the tube.
- g. **Add 5  $\mu$ L of the Negative Control nucleic acid to the last SmartCycler tube containing RT-PCR Master mix.**
  - i. After adding the Negative Control nucleic acid to the SmartCycler tube, **pipet up and down 2 to 3 times in the upper part of the tube.**
  - ii. Close the tube.
- h. **Centrifuge all tubes**
  - i. Appropriately label the SmartCycler tubes on the caps.
  - ii. Centrifuge all tubes for 5 to 10 seconds using the Cepheid microcentrifuge specially adapted to fit the SmartCycler tubes.
  - iii. Return tubes to the cooling block.
- i. **Keep the tubes on the Cooling Block before loading them into the SmartCycler instrument.**

### 5. Run the ProParaflu+ Assay (Amplification/Detection Area)

- a. Create a new run by clicking on the **Create Run** icon at the top of the screen. This will open the *Create Run* screen.
- b. Under **Run Name** in the left panel of the *Create Run* screen, enter a unique run identifier.
- c. Click on the **Assay** arrow in the left panel of the *Create Run* screen and select the **ProParaflu+ Assay** protocol from the drop-down menu.
- d. Under **Assay Information** in the left panel of the *Create Run* screen, enter the **Lot Number** and **Expiration Date** of the ProParaflu+ Supermix.
- e. In the left panel of the *Create Run* screen, enter the number of specimens (including the Positive and Extraction Controls but excluding the Negative Control.) and click on **Apply**. This will display the **Site Table** and the SmartCycler Dx Software will automatically select the **I-Core** sites.
- f. In the **Site Table** under the **Sample ID** column, enter the Sample Identifier or Positive Control Identifier for the appropriate I-Core sites.
- g. Insert each reaction tube into an I-Core site of the SmartCycler by pressing down firmly on all tubes and close each lid. Verify that the Negative Control (NC1) is loaded into the correct I-Core site.
- h. Select the **Start Run** button located at the bottom left corner of the screen. Verify that the LED is on for the appropriate I-Core sites.

### 6. Print report

- a. Click on **Report** at bottom of screen to open the **Report Preview** screen.
- b. Click on the **Print Icon** at the top of the screen.

**Interpretation of Control Results**
**Validation of Run**

**Note**

The user must interpret the Positive Control (PC) results and the Extraction Control (if included) results to determine whether the RT-PCR run and/or the extraction run are VALID; the SmartCycler Dx software will automatically interpret the Negative Control result.

For a **VALID RT-PCR run**, the following conditions must be met:

Sample ID <sup>1</sup>	Assay Result	IC Result	Warning / Error Code	Sample Type	IC Ct	HPIV-1 Result	HPIV-1 Ct	HPIV-3 Result	HPIV-3 Ct	HPIV-2 Result	HPIV-2 Ct
HPIV Cntrl <sup>2</sup>	Positive	NA	**	SPEC	NA	POS	20-40	POS	20-40	POS	20-40
Neg Cntrl	Valid <sup>3</sup>	Pass		NC1	15-50	Valid	0	Valid	0	Valid	0

<sup>1</sup> Columns and data not used for interpretation are not included.

<sup>2</sup> The Positive Control contains RNA for each of the Parainfluenza types (1 -3). The control needs to be valid/positive in all the channels, but the IC channel. Periodically bleedover may be observed in the IC (Cy5) channel, but this does not invalidate the run results.

<sup>3</sup> (Typical) an Invalid assay will display Error Code 4098.

\*\* Error Code 3079: Warning/Error Code 3079 is periodically observed with HPIV positives (Positive Control, Extraction Control, HPIV positive NP swab samples). Warning/Error Code 3079 occurs when the fluorescence (RFU) signal is too high. In this case, all results for the sample are reported by the Dx software as ND (Not Determined). When this code is observed for the Positive Control run validity can be determined based on Ct values of the Positive Control. The Positive Control must have a Ct value between 20-40 in the HPIV-1, HPIV-3 and HPIV-2 Ct columns to be considered VALID.

**Invalid RT-PCR Run**

If the Positive Control is not positive within the specified Ct range but the Negative Control is valid, prepare all new reactions using remaining purified nucleic acids and a new Positive Control (starting with PCR at [Step 4](#) of the [Assay Procedure](#)).

If the Negative Control is invalid, prepare all new extractions starting from original sample(s) using a new Extraction Control and a new Negative Control (starting at [Step 1](#) of the [Assay Procedure](#)).

For a **VALID Extraction run**, the following conditions must be met:

Sample ID <sup>1</sup>	Assay Result	IC Result	Warning / Error Code	Sample Type	IC Ct	HPIV-1 or HPIV-2 or HPIV-3 Result	HPIV-1 or HPIV-2 or HPIV-3 Ct
Extraction Control	Positive	NA	**	SPEC	NA	POS	20-40
Neg Cntrl	Valid <sup>2</sup>	Pass		NC1	15-50	Valid	0

<sup>1</sup> Columns and data not used for interpretation are not included.

<sup>2</sup> (Typical) an Invalid assay will display Error Code 4098.

\*\* Error Code 3079: Warning/Error Code 3079 is periodically observed with HPIV positives (Positive Control, Extraction Control, HPIV positive NP swab samples). Warning/Error Code 3079 occurs when the fluorescence (RFU) signal is too high. In this case, all results for the sample are reported by the Dx software as ND (Not Determined). When this code is observed for the Extraction Control, extraction run validity can be determined based on Ct values of the Extraction Control. The Extraction Control must have a Ct value between 20-40 in the HPIV-1 or HPIV-3 or HPIV-2 Ct column to be considered VALID.

**Invalid Extraction Run**

If the conditions for a valid extraction run are not met (i.e., the Extraction Control is not positive within the specified Ct range or the Negative Control is invalid), repeat the entire extraction run start from original sample(s) using a new Extraction Control and a new Negative Control (starting at [Step 1](#) of the [Assay Procedure](#)).

## Interpretation of Specimen Results

The SmartCycler Dx software automatically determines the specimen results. The interpretation of the assay specimen results is as follows:

Sample ID <sup>1</sup>	Assay Result	IC Result	Warning/Error Code	HPIV-1 Result	HPIV-3 Result	HPIV-2 Result	Interpretation of Results
Sample ID	Negative	Pass		<b>NEG</b>	<b>NEG</b>	<b>NEG</b>	HPIV-1, -2 and -3 nucleic acid <b>not detected</b>
Sample ID	Positive	NA*		<b>POS</b>	NEG	NEG	HPIV-1 nucleic acid <b>detected</b>
Sample ID	Positive	NA*		NEG	<b>POS</b>	NEG	HPIV-3 nucleic acid <b>detected</b>
Sample ID	Positive	NA*		NEG	NEG	<b>POS</b>	HPIV-2 nucleic acid <b>detected</b>
Sample ID	Positive	NA*		<b>POS</b>	<b>POS</b>	NEG	HPIV-1 and HPIV-3 nucleic acid <b>detected</b> – multiple infections are rare, repeat testing from purified nucleic acid or re-test from original sample. <b>Other combinations (HPIV-1 and HPIV-2 or HPIV-2 and HPIV-3) may occur in the same manner.</b>
Sample ID	Unresolved	Fail		<b>NEG</b>	<b>NEG</b>	<b>NEG</b>	Unresolved – PCR inhibition or reagent failure. Repeat testing from the purified nucleic acid or re-test from original sample.
Sample ID	ND <sup>2</sup>	ND	3079 <sup>2</sup>	ND	ND	ND	Not Determined – error code 3079
Sample ID	Invalid		4098 <sup>3</sup>	ND	ND	ND	Not Determined – error code 4098

<sup>1</sup> Columns and data not used for interpretation are not included

<sup>2</sup> Error Code 3079: Warning/Error Code 3079 is periodically observed. Warning/Error Code 3079 occurs when the fluorescence (RFU) signal is too high. In this case, all results for the sample are reported by the Dx software as ND (Not Determined). If a Ct value  $\geq 13$  is reported in any analyte column, results can be recorded as POSITIVE for that analyte.

<sup>3</sup> An Invalid Assay run will display Error code 4098

\* Detection of the Internal Control in the Cy5 detection channel is not required for positive result. High viral load can lead to reduced or absent Internal Control signal.

### Dual or Multiple Parainfluenza Infections

Dual or multiple parainfluenza infections are rare, and they may be artifacts of the SmartCycler Dx software due to signal bleed-over. It is required that repeat testing for these samples be performed starting from the purified nucleic acid, original sample, or a newly collected sample.

### Not Determined Samples

If an assay result of **ND** (Not Determined) is reported with an instrument failure other than Warning/Error Code 3079, repeat testing from the purified nucleic acids (starting with PCR, see [Step 4 \(a\)](#) of the [Assay Procedure](#)). Refer to the Cepheid Dx Software Operator Manual for interpretation of Warning Codes.

## Quality Control

- Quality control requirements must be performed in conformance with local, state, and/or federal regulations or accreditation requirements and your laboratory's standard quality control procedures. It is recommended that the user refer to CLSI document C24-A3, *Statistical Quality Control for Quantitative Measurements: Principles and Definitions*: [Approved Guideline – Third Edition] or other published guidelines for general quality control recommendations. For further guidance on appropriate quality control practices, refer to 42 CFR 493.1202(c).
- Quality control procedures are intended to monitor reagent and assay performance.

Control Type	Used to Monitor
Positive	Substantial reagent failure including primer and probe integrity
Negative	Reagent and/or environmental contamination
Extraction	Failure in lysis and extraction procedure
Internal	PCR inhibition in individual samples and Reagent failure or process error

- Dilute and test the Positive Control and the Internal Control prior to running samples with each new kit lot to ensure all reagents and kit components are working properly.
- Good laboratory practice recommends including a positive Extraction Control (not provided) in each nucleic acid isolation run. The Extraction Control should be treated as a sample.
- Never run the Positive Control through nucleic acid isolation.
- Always include a Negative Control (*containing Internal Control*) and the Positive Control in each amplification/detection run performed.
- Failure of Controls (Positive, Negative and/or Extraction) invalidates the run and results should not be reported.
- If the Positive Control is not positive within the specified Ct range but the Negative Control is valid, repeat testing should be done starting from the purified nucleic acid and using a new aliquot of the Positive Control. If repeat results are still invalid, results should not be reported and testing should be repeated from the original sample or a new sample should be collected and tested.
- If the Extraction Control is not positive within the specified Ct range or the Negative Control is invalid, repeat testing should be done starting from the original sample and using a new Extraction Control and a new Negative Control. If repeat results are still invalid, results should not be reported and a new sample should be collected and tested.

## Limitations

- **Once the RT-PCR Master mix has been made, the run must be started within one hour.**
- This test does not detect Parainfluenza 4a or Parainfluenza 4b.
- Negative test results are presumptive and should be confirmed by cell culture. Negative results do not preclude Parainfluenza 1, 2 or 3 virus infections and should not be used as the sole basis for treatment or other management decisions.
- Dual or multiple parainfluenza infections are rare, and they may be artifacts of the SmartCycler Dx software due to signal bleed-over. It is required that repeat testing for these samples be performed starting from the purified nucleic acid, original sample, or a newly collected sample.
- Optimal assay performance requires strict adherence to the assay procedure describe in this insert.
- A trained health care professional should interpret assay results in conjunction with the patient's medical history, clinical signs and symptoms, and the results of other diagnostic tests.
- Analyte targets (viral nucleic acid) may persist *in vivo*, independent of virus viability. Detection of analyte target(s) does not imply that the corresponding virus(es) are infectious, or are the causative agents for clinical symptoms.
- The detection of viral nucleic acid is dependent upon proper specimen collection, handling, transportation, storage, and preparation (including extraction). Failure to observe proper procedures in any one of these steps can lead to incorrect results.
- There is a risk of false negative values resulting from improperly collected, transported, or handled specimens.
- There is a risk of false negative values due to the presence of sequence variants in the viral targets of the assay, procedural errors, amplification inhibitors in specimens, or inadequate numbers of organisms for amplification.
- A specimen yielding a negative result may contain respiratory viruses other than Parainfluenza 1, Parainfluenza 2 or Parainfluenza 3. A negative result should not be used as the sole basis for diagnosis, treatment or other management decisions.
- There is a risk of false positive values resulting from cross-contamination by target organisms, their nucleic acids or amplified product, or from non-specific signals in the assay.
- False negative results may occur due to loss of nucleic acid. The Internal Control has been added to the test to aid in the identification of specimens that contain inhibitors to PCR amplification. The Internal Control does not indicate whether or not nucleic acid has been lost due to inadequate collection, transport or storage of specimens.
- The ProParaflu+ Assay may not generate reproducibly positive results when testing samples that have analyte concentrations lower than the LoD concentration, but higher than the assay cutoff concentration.
- The performance of the ProParaflu+ Assay has not been established in immunocompromised patients.
- Positive and negative predictive values are highly dependent on prevalence. The assay performance was established during May 2008 to September 2009. The performance may vary depending on the prevalence and population tested.
- Some evidence of interference of HPIV detection at the LoD concentration was observed with throat lozenges, Relenza and Rebitol in the Interference Study, albeit these substances were tested at concentrations much higher than would be encountered *in vivo*.

**Expected Values**

In the US and around the world, infections with HPIV-1 and HPIV-2 occur in biennial alternating outbreaks during fall/early-winter where as HPIV-3 occurs throughout the year but generally peaks in spring<sup>6,7</sup>. Variables that affect the rate of positivity observed in respiratory testing include: the efficiency and timing of specimen collection, handling and transport of the specimen, the time of year, age of the patient, and local disease prevalence.

In the prospective ProParaflu+ Assay clinical study, a total of 857 eligible nasopharyngeal (NP) swab specimens were tested from four U.S. clinical laboratories across the United States from May 2008 to September 2009. The number and percentage of HPIV-1, HPIV-2 and HPIV-3 RNA positive cases as determined by the ProParaflu+ Assay, calculated by age group, are presented in the following table:

Age Group	Total N	Number HPIV-1 Positive By the ProParaflu+ Assay	Number HPIV-2 Positive By the ProParaflu+ Assay	Number HPIV-3 Positive By the ProParaflu+ Assay	Observed Prevalence HPIV-1	Observed Prevalence HPIV-2	Observed Prevalence HPIV-3
< 1 year	331	7	14	29	2.1%	4.2%	8.8%
1-5 years	249	9	9	11	3.6%	3.6%	4.4%
6-10 years	58	1	1	1	1.7%	1.7%	1.7%
11-15 years	61	0	1	0	0%	1.6%	0%
16-21 years	49	0	2	0	0%	4.1%	0%
> 21 years	109	0	1	3	0%	0.9%	2.8%
Total	857	17	28	43	2.0%	3.3%	5.0%

**Performance Characteristics**
**Clinical Performance**

The clinical performance of the ProParaflu+ Assay was established during a prospective study at 4 U.S. clinical laboratories during May 2008 – September 2009. Specimens used in the study represented excess nasopharyngeal (NP) swab specimens that were prospectively collected from symptomatic individuals suspected of respiratory infection, and were submitted for routine analysis. Demographic details for this patient population are summarized in the following table.

Gender and Age Demographic Detail for ProParaflu+ Prospective Study

Sex	Number of Subjects
Female	407 (47.5%)
Male	450 (52.5%)
Age (yrs)	
≤ 5 years	580 (67.7%)
6 - 21 years	168 (19.6%)
22 – 59 years	67 (7.8%)
≥ 60 years	42 (4.9%)

Performance of the ProParaflu+ Assay was compared to the reference method of cell culture (rapid or traditional) followed by direct fluorescent antibody (DFA) screening and HPIV type identification.

A total of 857 eligible NP swab samples were tested with the ProParaflu+ Assay and by culture across four clinical sites. Of the ProParaflu+ Assay run on all eligible specimens, 99.2% (852/857) of these specimens were successful on the first attempt. The remaining 5 gave “Unresolved” results on the first attempt. Unresolved results occur when the sample is negative for all three HPIVs and the Internal Control, indicating potentially PCR-inhibiting samples. Of the 5 “Unresolved” specimens on the first attempt, 60.0% (3/5) gave a valid result on the second attempt. The remaining 2 were “Unresolved” on the second attempt and are not included in the analysis below. Both samples were culture negative.

Discrepant analysis for samples where ProParaflu+ Assay and culture results were in disagreement was performed using RT-PCR with virus specific primers obtained from literature<sup>8,9</sup> (and different from those used in ProParaflu+) followed by bi-directional sequencing.

**Prospective Study**
**Parainfluenza 1 Comparison Results**

		Culture/DFA		Total	
		Positive	Negative		
<b>ProParaflu+ Assay</b>	<b>Positive</b>	16	1 <sup>a</sup>	<b>17</b>	Sensitivity 88.9% (67.2% - 96.9%) 95% CI
	<b>Negative</b>	2 <sup>b</sup>	838	<b>840</b>	Specificity 99.9% (99.3% - 100.0%) 95% CI
	<b>Total</b>	<b>18</b>	<b>839</b>	<b>857</b>	

<sup>a</sup>One (1) sample positive for HPIV-1 by bi-directional sequence analysis.

<sup>b</sup>Two (2) samples negative for HPIV-1 by bi-directional sequence analysis. One sample positive for HPIV-3 by ProParaflu+ and bi-directional sequence analysis.

**Parainfluenza 2 Comparison Results**

		Culture/DFA		Total	
		Positive	Negative		
ProParaflu+ Assay	Positive	26	2 <sup>a</sup>	<b>28</b>	Sensitivity 96.3% (81.7% - 99.3%) 95% CI
	Negative	1 <sup>b</sup>	828	<b>829</b>	Specificity 99.8% (99.1% - 99.9%) 95% CI
<b>Total</b>		<b>27</b>	<b>830</b>	<b>857</b>	

<sup>a</sup>Two (2) samples positive for HPIV-2 by bi-directional sequence analysis.

<sup>b</sup>One (1) sample negative for HPIV-2 by bi-directional sequence analysis.

**Parainfluenza 3 Comparison Results**

		Culture/DFA		Total	
		Positive	Negative		
ProParaflu+ Assay	Positive	36	8 <sup>a</sup>	<b>44</b>	Sensitivity 97.3% (86.2% - 99.5%) 95% CI
	Negative	1 <sup>b</sup>	812	<b>813</b>	Specificity 99.2% (98.1% - 99.5%) 95% CI
<b>Total</b>		<b>37</b>	<b>820</b>	<b>857</b>	

<sup>a</sup>Seven (7) samples positive for HPIV-3 and one (1) sample negative for HPIV-3 by bi-directional sequence analysis.

<sup>b</sup>One (1) sample negative for HPIV-3 by bi-directional sequence analysis.

**Retrospective Study**

Due to a minimal number of HPIV-1 positive samples, a retrospective study was also conducted using a total of 91 frozen NP swab samples that had been previously tested by direct DFA. Demographic details for this patient population are summarized in the following table.

Gender and Age Demographic Detail for ProParaflu+ Retrospective Study

Sex	Number of Subjects
Female	40 (44.4%)
Male	50 (55.6%)
Age (yrs)	
≤ 5 years	81 (90.0%)
6 - 21 years	5 (5.6%)
22 – 59 years	2 (2.2%)
≥ 60 years	2 (2.2%)

**Parainfluenza 1 Comparison Results**

		DFA		Total	
		Positive	Negative		
ProParaflu+ Assay	Positive	24	0	<b>24</b>	Sensitivity 82.8% (65.4% - 92.4%) 95% CI
	Negative	5 <sup>a</sup>	62	<b>67</b>	Specificity 100% (94.2% - 100%) 95% CI
<b>Total</b>		<b>29</b>	<b>62</b>	<b>91</b>	

<sup>a</sup>Five (5) samples negative for HPIV-1 by bi-directional sequence analysis.

### Reproducibility

The reproducibility of the ProParaflu+ Assay was evaluated at 3 laboratory sites. Reproducibility was assessed using a panel of 9 simulated samples that included medium positive, low positive (near the assay limit of detection,  $\geq 95\%$  positive), and high negative (below the assay limit of detection,  $< 5\%$  positive) samples. Panels and controls were tested at each site by 2 operators for 5 days (9 samples and 3 controls/run X 1 run/day/operator X 2 operators X 5 days X 3 sites = 360). Nucleic acid extraction on the test panel samples were carried out using either the Roche MagNA Pure LC System (Site #3) or the bioMérieux NucliSENS easyMAG System (Site #1 and Site #2). The overall percent agreement with the expected result for the ProParaflu+ Assay was 97.8%.

	Panel Member ID	HPIV-1 high negative <sup>a</sup>	HPIV-1 low positive	HPIV-1 medium positive	HPIV-2 high negative <sup>a</sup>	HPIV-2 low positive	HPIV-2 medium positive	HPIV-3 high negative <sup>a</sup>	HPIV-3 low positive	HPIV-3 medium positive	Para Extraction Control	Para RNA Control			Negative Control <sup>a</sup>	Total % Agreement
		Concentration	HPIV-1	HPIV-2	HPIV-3	HPIV-1	HPIV-2	HPIV-3								
	Concentration	0.001 X LoD	2 X LoD	10X LoD	0.001 X LoD	2 X LoD	10X LoD	0.01 X LoD	2 X LoD	10X LoD	N/A	N/A			N/A	
Site 1	Agreement with Expected Result	10/10 100%	8/10 80%	9/9 100%	10/10 100%	9/9 100%	9/10 90%	10/10 100%	9/10 90%	10/10 100%	10/10 100%	10/10 100%			10/10 100%	<b>114/118</b> <b>96.6%</b>
	Mean Ct Value	27.73	28.31	26.33	27.90	28.62	26.27	27.76	31.21	29.47	27.33	27.37	29.37	28.61	27.69	
	% CV	2.87	1.55	1.60	2.84	0.85	1.25	2.43	3.21	1.91	1.60	1.05	0.43	0.81	1.67	
Site 2	Agreement with Expected Result	8/10 80%	8/10 80%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%			10/10 100%	<b>116/120</b> <b>96.7%</b>
	Mean Ct Value	28.59	28.47	26.12	28.83	28.91	26.61	28.30	31.64	29.51	27.56	23.86	26.09	25.23	28.68	
	% CV	1.36	1.72	1.26	2.80	1.31	1.83	1.18	2.17	2.66	2.47	1.48	1.12	0.92	1.15	
Site 3	Agreement with Expected Result	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%	10/10 100%			10/10 100%	<b>120/120</b> <b>100%</b>
	Mean Ct Value	26.35	29.91	27.67	26.28	29.51	27.44	26.67	33.13	30.43	28.61	28.73	30.98	29.84	26.46	
	% CV	0.88	0.81	1.25	1.35	0.57	2.25	4.13	2.36	1.02	1.54	3.31	3.39	3.19	1.04	
	Total Agreement with Expected Result	28/30 93.3%	26/30 86.7%	29/29 100%	30/30 100%	29/29 100%	29/30 96.7%	30/30 100%	29/30 96.7%	30/30 100%	30/30 100%	30/30 100%			30/30 100%	<b>350/358</b> <b>97.8%</b>
	95% CI	78.7% - 98.2%	70.3% - 94.7%	88.3% - 100%	88.6% - 100%	88.3% - 100%	83.3% - 99.4%	88.6% - 100%	83.3% - 99.4%	88.6% - 100%	88.6% - 100%	88.6% - 100%			88.6% - 100%	
	Overall Mean Ct Value	27.56	28.90	26.72	27.67	29.03	26.79	27.58	32.02	29.80	27.83	26.65	28.81	27.89	27.62	
	Overall % CV	3.88	2.87	2.95	4.55	1.57	2.54	3.68	3.61	2.42	2.75	8.13	7.49	7.39	3.54	

<sup>a</sup>Average Ct value for the Internal Control (IC)

An additional reproducibility study was performed to assess samples that were at an intermediate concentration, below the assay's LoD but above the "high negatives" tested during the original reproducibility study. Panels and controls were tested at each site by 2 operators for 5 days (3 samples and 3 controls/run X 1 run/day/operator X 2 operators X 5 days X 3 sites = 180). Nucleic acid extraction on the test panel samples were carried out using either the Roche MagNA Pure LC System (Site #3) or the bioMérieux NucliSENS easyMAG System (Site #1 and Site #2). The percent positive for the intermediate member across all sites was 56.7% for HPIV-1 (mean Ct = 35.1), 86.7% for HPIV-2 (mean Ct = 33.0), and 30.0% for HPIV-3 (mean Ct = 37.1). This result was expected as the intermediate concentration should be positive in the range of 5 - 95% as the samples were lower concentration than the LoD concentration ( $\geq 95\%$  positive) and higher than the "high negative" concentration ( $< 5\%$  positive).

Panel Member ID	HPIV-1 Intermediate	HPIV-2 Intermediate	HPIV-3 Intermediate	Para Extraction Control	Parainfluenza RNA Control			Negative Control <sup>a</sup>	
					HPIV-1	HPIV-2	HPIV-3		
Concentration	0.1 X LoD	0.1 X LoD	0.1 X LoD	N/A	N/A			N/A	
Site 1	Agreement with Positive Result	4/10 40%	8/10 80%	1/10 10%	10/10 100%	10/10 100%			10/10* 100%
	Average Ct Value	35.5	33.3	36.9	27.9	28.8	30.4	29.5	28.4
	% CV	6.19	2.78	N/A	3.80	1.13	0.88	0.89	3.47
Site 2	Agreement with Positive Result	8/10 80%	10/10 100%	7/10 70%	10/10 100%	10/10 100%			10/10 100%
	Average Ct Value	34.4	32.2	37.3	27.5	29.0	30.6	29.9	27.7
	% CV	1.38	2.22	1.50	2.92	1.04	0.89	0.72	2.80
Site 3	Agreement with Positive Result	5/10 50%	8/10 80%	1/10 10%	10/10 100%	10/10 100%			10/10 100%
	Average Ct Value	35.9	33.7	35.8	28.6	29.7	31.5	30.3	27.9
	% CV	2.44	2.90	N/A	2.75	1.92	1.69	1.46	1.94
Total Agreement with Positive Result	17/30 56.7%	26/30 86.7%	9/30 30.0%	30/30 100%	30/30 100%			30/30 100%	
95% CI	39.2% - 72.6%	70.3% - 94.7%	16.7% - 47.9%	88.7 - 100%	88.7 - 100%			88.7 - 100%	
Overall Average Ct Value	35.1	33.0	37.1	28.0	29.1	30.8	29.9	28.0	
Overall % CV	3.68	3.25	1.89	3.46	1.91	1.92	1.53	2.92	

<sup>a</sup>Average Ct value for the Internal Control (IC)

\*Agreement with Negative result

### Analytical Sensitivity

The analytical sensitivity (limit of detection or LoD) of the ProParaflu+ Assay was determined using quantified (TCID<sub>50</sub>/mL) cultures of HPIV-1, HPIV-2 and HPIV-3 serially diluted in nasopharyngeal clinical matrix. Each viral strain was extracted using the Roche MagNA Pure LC instrument and tested in replicates of 20 per concentration of virus.

Analytical sensitivity (LoD), defined as the lowest concentration at which  $\geq 95\%$  of all replicates tested positive, ranged from  $5 \times 10^1 - 10^2$  TCID<sub>50</sub>/mL.

Virus	Strain	LoD Concentration
HPIV-1	C35	$10^2$ TCID <sub>50</sub> /mL
HPIV-2	Greer	$10^2$ TCID <sub>50</sub> /mL
HPIV-3	C 243	$5 \times 10^1$ TCID <sub>50</sub> /mL

### Analytical Specificity

The analytical specificity of the ProParaflu+ Assay was evaluated by testing a panel of 27 viruses, 24 bacteria, and 1 yeast strain representing common respiratory pathogens or flora commonly present in nasopharynx. Bacteria and yeast were tested at concentrations of  $10^6$  to  $10^8$  CFU/mL. Viruses were tested at concentrations of  $10^3$  to  $10^6$  TCID<sub>50</sub>/mL. HPIV-1, 2 and 3 viruses were tested near and above LoD. Samples were extracted using the Roche MagNA Pure LC instrument and tested in triplicate. Analytical specificity of the ProParaflu+ Assay was 100%.

Strains	Concentration	HPIV-1 (FAM)	HPIV-3 (TET)	HPIV-3 (Tex Red)
Parainfluenza Type 1	10 <sup>4</sup> TCID <sub>50</sub> /mL	+	-	-
Parainfluenza Type 1	5x10 <sup>2</sup> TCID <sub>50</sub> /mL	+	-	-
Parainfluenza Type 2	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	+
Parainfluenza Type 2	5x10 <sup>2</sup> TCID <sub>50</sub> /mL	-	-	+
Parainfluenza Type 3	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	+	-
Parainfluenza Type 3	5x10 <sup>1</sup> TCID <sub>50</sub> /mL	-	+	-
Parainfluenza Type 4	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Adenovirus 1/Adenoid 71	10 <sup>6</sup> TCID <sub>50</sub> /mL	-	-	-
Coronavirus 229E	10 <sup>6</sup> TCID <sub>50</sub> /mL	-	-	-
Coxsackie B4	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Coxsackie B5/10/2006	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	-	-
Cytomegalovirus	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Echovirus 2	10 <sup>6</sup> TCID <sub>50</sub> /mL	-	-	-
Echovirus 3	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	-	-
Echovirus 6	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	-	-
Echovirus 11	10 <sup>6</sup> TCID <sub>50</sub> /mL	-	-	-
Enterovirus 68	10 <sup>3</sup> TCID <sub>50</sub> /mL	-	-	-
Enterovirus 70	10 <sup>3</sup> TCID <sub>50</sub> /mL	-	-	-
hMPV A2	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
HSV Type 1 MacIntyre Strain	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	-	-
HSV Type 2 G strain	10 <sup>5</sup> TCID <sub>50</sub> /mL	-	-	-
Human Rhinovirus 39	10 <sup>3</sup> TCID <sub>50</sub> /mL	-	-	-
Human Rhinovirus	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Influenza A/Port Chalmers	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Influenza B/Wisconsin	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Measles/7/2000	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Mumps Virus	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
RSV A Strain Long	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
RSV B Strain Wash	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
Varicella Zoster Virus	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
<i>Bordetella pertussis</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Bordetella bronchiseptica</i>	5x10 <sup>7</sup> CFU/mL	-	-	-
<i>Chlamydia pneumoniae</i>	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
<i>Chlamydia trachomatis</i>	10 <sup>4</sup> TCID <sub>50</sub> /mL	-	-	-
<i>Legionella pneumophila</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Mycobacterium intracellulare</i>	10 <sup>7</sup> CFU/mL	-	-	-
<i>Mycobacterium tuberculosis</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Haemophilus influenzae</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Pseudomonas aeruginosa</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Proteus vulgaris</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Proteus mirabilis</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Neisseria gonorrhoeae</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Neisseria meningitidis</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Neisseria mucosa</i>	7.4x 10 <sup>7</sup> CFU/mL	-	-	-
<i>Klebsiella pneumoniae</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Escherichia coli</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Moraxella catarrhalis</i>	1.3x10 <sup>7</sup> CFU/mL	-	-	-
<i>Corynebacterium diphtheriae</i>	3x10 <sup>7</sup> CFU/mL	-	-	-
<i>Lactobacillus plantarum</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Streptococcus pneumoniae</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Streptococcus pyogenes</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Streptococcus salivarius</i>	2x10 <sup>5</sup> CFU/mL	-	-	-
<i>Staphylococcus epidermidis</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Staphylococcus aureus</i>	10 <sup>6</sup> CFU/mL	-	-	-
<i>Candida albicans</i>	10 <sup>6</sup> CFU/mL	-	-	-

Note: Fresh cultured and titered stocks of organisms were used for the analytical specificity study, except for *C. pneumoniae* and *C. trachomatis*. ATCC frozen cultures and ATCC supplied titers were used for both due to technical difficulties in re-growing and re-titering these organisms.

### Interference

Mucin, whole blood and a number of potentially interfering exogenous substances (medications and over the counter (OTC) products) that may be present in the nasopharynx were evaluated in the ProParaflu+ Assay. HPIV-3 was spiked into HPIV negative NP pools at 2X LoD and 10X LoD. Clinically relevant amounts of the potential interfering substances were added to spiked samples. An Internal Control (IC) was also added to each sample. Nucleic acid from the samples was extracted with the Roche MagNA Pure LC instrument. The ProParaflu+ Assay was performed in triplicate reactions for each sample on the Cepheid SmartCycler II. The following table shows the potential interfering substances used for this study. The substances consisted of nasal sprays (liquid and powder), ingestible pills and lozenges, injectables, and endogenous substances:

Substance Name	Active Ingredient	Concentration Tested
Mucin	Purified mucin protein	60 µg/mL
Blood (human)	N/A	2% (volume/volume)
Neo-Syneprine	Phenylephrine HCl	15% (volume/volume)
Anefrin Nasal Spray	Oxymetazoline Hydrochloride	15% (volume/volume)
Zicam Nasal gel	Luffa Operculata, Galphimia Glauca, Histaminum Hydrochloricum,	5% (volume/volume)
Saline Nasal Spray	Sodium chloride with preservatives	15% (volume/volume) of dose
Chloraseptic Throat Lozenges	Oral anesthetic/analgesic	0.63 mg/mL; active ingredients: 1.0 mg/mL benzocaine, 1.7 mg/mL menthol
Relenza	Zanamivir	3.3 mg/mL
Tobramycin	Tobramycin	4.0 µg/mL
Mupirocin	Mupirocin	6.6 mg/mL
Rebitol	Ribavirin	20 mg/mL
TamiFlu	Oseltamivir	25 mg/mL
Beconase AQ	Beclomethasone dipropionate	5% (volume/volume)

All of the exogenous or endogenous potentially interfering substances tested were found to have no effect on ProParaflu+ Assay results when present in simulated respiratory samples at the 10X LoD concentration. However, some evidence of interference of HPIV detection at the LoD was observed with throat lozenges, Relenza and Rebitol, albeit these substances were tested at concentrations much higher than would be encountered *in vivo*.

### Extraction Equivalency

Extraction equivalency of the bioMérieux NucliSENS easyMAG and Roche MagNA Pure LC instruments was evaluated by performing a limit of detection study. A single cultured and titered strain of HPIV-3 was spiked (along with an IC) into individual aliquots of negative NP matrix pools at concentrations of 1 log above, at, and 1 log below the previously determined LoD. Each viral strain dilution was extracted in replicates of 10 on each automated extractor and tested using the ProParaflu+ Assay.

The bioMérieux NucliSens easyMAG instrument and the Roche MagNA Pure LC instrument performed equivalently with respect to limit of detection.

### Carry-over/Cross-Contamination

To evaluate the degree of carry-over/cross-contamination that occurs with the use of the ProParaflu+ Assay in association with nucleic acid extraction on the Roche MagNA Pure LC and the bioMérieux NucliSens easyMAG instruments and RT-PCR on the Cepheid SmartCycler II thermocycler, an internal Carry-Over study was performed. Simulated HPIV-2 high positive samples were run in series alternating with HPIV-2 high negative samples. The HPIV-2 high positive samples in this study represented the lower Cycle Threshold (Ct) range (higher sample titer range) obtained in the Pro hMPV+ clinical trials (lowest Ct = 22.8). The high-negative samples include a low amount of HPIV-2 that should be detectable no more than 5% of the time. Five (5) out of 110 (4.5%) high-negative samples tested showed potential HPIV-2 contamination when extracted using either extraction system in the study. Also some HPIV-1 and HPIV-3 contamination was also observed (n = 5 of 220) during initial testing, but was not observed when samples were retested. Potential contamination could have occurred during: creation of the sample, sample preparation for extraction, extraction, transfer of purified nucleic acid samples from the sample cartridge to microfuge tubes, or set up of the RT-PCR reaction. The possibility that these high negative samples (HPIV-2) fall in the 5% category could not be ruled out.

### Disposal

Dispose of hazardous or biologically contaminated materials according to the practices of your institution.

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








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