

AdV/hMPV/RV Assay (Panther Fusion™ System)

For in vitro diagnostic use.

For U.S. Export Only.

CONTENTS

General Information	2
Intended Use	2
Summary and Explanation of the Test	2
Principles of the Procedure	3
Warnings and Precautions	4
Reagent Storage and Handling Requirements	ô
Specimen Collection and Storage	7
Specimen Transport	8
Panther Fusion System	9
Reagents and Materials Provided for Panther Fusion AdV/hMPV/RV Assay	9
Materials Required and Available Separately	0
Panther Fusion System Test Procedure	1
Procedural Notes	2
Quality Control	2
Interpretation of Results	3
Limitations	4
Panther Fusion System Assay Performance	5
Clinical Performance	5
Analytical Sensitivity	ô
Reactivity17	7
Analytical Specificity	9
Competitive Interference	1
Interference	1
Carry-Over/Contamination	2
Assay Precision	3
Bibliography 25	5

General Information

Intended Use

The Panther Fusion™ AdV/hMPV/RV assay is a multiplex real-time PCR (RT-PCR) *in vitro* diagnostic test for the rapid and qualitative detection and differentiation of Adenovirus (AdV), human Metapneumovirus (hMPV), and Rhinovirus (RV). Nucleic acids are isolated and purified from nasopharyngeal (NP) swab specimens obtained from individuals exhibiting signs and symptoms of a respiratory tract infection.

This assay is intended to aid in the differential diagnosis of Adenovirus, human Metapneumovirus, and Rhinovirus infections in humans. Negative results do not preclude Adenovirus, human Metapneumovirus, and Rhinovirus infections and should not be used as the sole basis for treatment or other management decisions. This assay is designed for use on the Panther Fusion system.

Summary and Explanation of the Test

Respiratory viruses are responsible for a wide range of acute respiratory tract infections including the common cold, influenza, and croup and represent the most common cause of acute illness in the United States. Disease severity can be especially high in the young, the immunocompromised, and elderly patients. Accurate and timely diagnosis of the cause of respiratory tract infections has many benefits. They include improved treatment of the patient by ensuring appropriate antiviral treatment (e.g. oseltamivir for influenza), decreasing the overall cost of care, reducing selection for antimicrobial resistant organisms due to excessive and inappropriate use of antibiotics,¹ assisting infection control personnel in providing appropriate measures to minimize nosocomial spread, and providing valued information to public health authorities regarding which viruses are circulating in the community.²

Adenoviruses are members of the *Adenoviridae* family which are medium-sized (90-100 nm), non-enveloped icosohedral viruses with double-stranded DNA.³ At this time, in humans there are over 50 Adenovirus types in seven species (A to G).⁴ Adenoviruses most commonly cause respiratory illness which can range from the common cold to pneumonia, croup, and bronchitis.³ Depending on the type, Adenoviruses can cause other illnesses such as gastroenteritis, conjunctivitis, cystitis, and, less commonly, neurological disease.³ Infants and people with weakened immune systems are at high risk for developing severe illness caused by Adenovirus infection.³ Adenovirus circulates year-round and outbreaks are more common in late winter, spring, and early summer but can occur throughout the year.⁵

Since the discovery of hMPV in 2001, the virus has been identified worldwide. hMPV is a common respiratory pathogen, particularly in infants and young children. The virus is associated with both upper and lower respiratory tract infections and may be a trigger for asthma.⁶ Symptoms commonly associated with hMPV include cough, fever, nasal congestion, and shortness of breath. Clinical symptoms of hMPV infection may progress to bronchiolitis or pneumonia and are similar to other viruses that cause upper and lower respiratory infections. The incubation period is estimated to be 3 to 6 days, and the median duration of illness can vary depending upon severity but is similar to other respiratory infections caused by viruses.⁷ The peak of incidence of hMPV is mainly in the spring in temperate latitudes.⁸

Rhinoviruses, members of the family Picornaviridae, are the causative pathogens in more than half of viral respiratory infections, and they are associated with acute exacerbations of

respiratory disease, including asthma, sinusitis, otitis media, and COPD.9 A number of studies have confirmed rhinoviruses as being the most common cause of "the common cold" and affect all age groups.8 Symptoms usually include sore throat, runny nose, coughing, sneezing, watery eyes, headaches and body aches. Most people recover within about 7-10 days.8 Rhinoviruses circulate year round and tend to peak in the spring and fall.8

Principles of the Procedure

The Panther Fusion AdV/hMPV/RV assay involves the following steps: sample lysis, nucleic acid capture and elution transfer, and multiplex RT-PCR when analytes are simultaneously amplified, detected and differentiated. Nucleic acid capture and elution takes place in a single tube on the Panther Fusion system. The eluate is transferred to the Panther Fusion system reaction tube containing the assay reagents. Multiplex RT-PCR is then performed for the eluted nucleic acid on the Panther Fusion system.

Nucleic acid capture and elution: Prior to processing and testing on the Panther Fusion system, specimens are transferred to a Specimen Lysis Tube containing specimen transport media (STM) that lyses the viral particles, releases target nucleic acid and protects it from degradation during storage.

The Internal Control-S (IC-S) is added to each test specimen and controls via the working Panther Fusion Capture Reagent-S (wFCR-S). The IC-S in the reagent monitors specimen processing, amplification and detection.

Capture oligonucleotides hybridize to nucleic acid in the test specimen. Hybridized nucleic acid is then separated from the specimen in a magnetic field.

Wash steps remove extraneous components from the reaction tube. The elution step elutes purified nucleic acid. During the nucleic acid capture and elution step, total nucleic acid is isolated from specimens.

Elution transfer and RT-PCR: During the elution transfer step, eluted nucleic acid is transferred to a Panther Fusion reaction tube already containing oil and reconstituted mastermix.

For RV, hMPV, and internal control targets, amplification occurs via RT-PCR. A reverse transcriptase step generates DNA copies of the target sequence. For AdV, target amplification occurs via PCR. For all targets, specific forward and reverse primers and probes amplify targets while simultaneously detecting and discriminating multiple target types via multiplex PCR.

The Panther Fusion system compares the fluorescence signal to a predetermined cut-off to produce a qualitative result for the presence or absence of the analyte.

The analytes and the channel used for their detection on the Panther Fusion system is summarized in the table below.

Analyte	Gene Targeted	Instrument Channel
Adenovirus	Hexon	HEX
human Metapneumovirus	Nucleocapsid	ROX
Rhinovirus	5' UTR	FAM
Internal Control	Not applicable	RED677

Warnings and Precautions

- A. For *in vitro* diagnostic use.
- B. Carefully read this entire package insert and the Panther Fusion System Operator's Manual.
- C. The Panther Fusion Enhancer Reagent-S (FER-S) is corrosive, harmful if swallowed and causes severe skin burns and eye damage.
- D. Only personnel adequately trained on the use of this assay and in handling potentially infectious materials should perform these procedures. If a spill occurs, immediately disinfect using appropriate site procedures.
- E. 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.
- F. Use only supplied or specified disposable laboratory ware.
- G. Wear disposable, powderless gloves, protective eye wear, and laboratory coats when handling specimens and reagents. Wash hands thoroughly after handling specimens and reagents.
- H. Dispose of all material that has come into contact with specimens and reagents in accordance with applicable national, international, and regional regulations.
- I. Expiration dates listed on the Panther Fusion Specimen Lysis Tubes pertain to the transfer of sample into the tube and not to testing of the sample. Specimens collected/transferred any time prior to these expiration dates are valid for testing provided they are transported and stored in accordance with the appropriate package insert, even if these expiration dates have passed.
- J. Maintain proper storage conditions during specimen shipping to ensure the integrity of the specimen. Specimen stability under shipping conditions other than those recommended has not been evaluated.
- K. Avoid cross-contamination during the specimen handling steps. Specimens can contain extremely high levels of virus or other organisms. Ensure that specimen containers do not come in contact with one another, and discard used materials without passing them over any open containers. Change gloves if they come in contact with specimens.
- L. Do not use the reagents and controls after the expiration date.
- M. Store assay components at the recommended storage condition. See *Reagent Storage and Handling Requirements* (page 6), and *Panther Fusion System Test Procedure* (page 11) for more information.
- N. Do not combine any assay reagents or fluids. Do not top off reagents or fluids; the Panther Fusion system verifies reagent levels.
- O. Avoid microbial and ribonuclease contamination of reagents.

- P. 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. Reference 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 is recommended. For further guidance on appropriate quality control practices, refer to 42 CFR 493.1205.
- Q. Do not use the assay cartridge if the storage pouch has lost its seal or if the assay cartridge foil is not intact. Contact Hologic if either occurs.
- R. Do not use the fluid packs if the foil seal is leaking. Contact Hologic if this occurs.
- S. Handle the assay cartridges with care. Do not drop or invert assay cartridges. Avoid prolonged exposure to ambient light.



Panther Fusion Oil Polydimethylsiloxane 100%

Warning

H315 - Causes skin irritation

H319 - Causes serious eye irritation



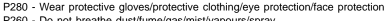
Panther Fusion Enhancer Reagent-S Lithium Hydorxide Monohydrate 5-10%



Danger

H302 - Harmful if swallowed





P260 - Do not breathe dust/fume/gas/mist/vapours/spray P303 + P361 + P353 - IF ON SKIN (or hair): Remove/Take off immediately all contaminated clothing. Rinse skin with

water/shower P280 - Wear eye protection/ face protection

P305 + P351 + P338 - IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing

P310 - Immediately call a POISON CENTER or doctor/physician

Note: For information on any hazard and precautionary statements that may be associated with reagents, refer to the Safety Data Sheet Library at www.hologic.com/sds.

Reagent Storage and Handling Requirements

A. The following table provides storage and handling requirements for this assay.

Reagent	Unopened Storage	On Board/ Open Stability ¹	Opened Storage
Panther Fusion AdV/hMPV/RV Assay Cartridge	2°C to 8°C	60 days	2°C to 8°C ²
Panther Fusion Capture Reagent-S (FCR-S)	15°C to 30°C	30 days	15°C to 30°C
Panther Fusion Enhancer Reagent-S (FER-S)	15°C to 30°C	30 days	15°C to 30°C
Panther Fusion Internal Control-S (IC-S)	2°C to 8°C	(In wFCR-S)	Not applicable
Panther Fusion Elution Buffer	15°C to 30°C	60 days	15°C to 30°C
Panther Fusion Oil	15°C to 30°C	60 days	15°C to 30°C
Panther Fusion Reconstitution Buffer I	15°C to 30°C	60 days	15°C to 30°C
Panther Fusion AdV/hMPV/RV Positive Control	2°C to 8°C	Single use vial	Not applicable- single use
Panther Fusion Negative Control	2°C to 8°C	Single use vial	Not applicable- single use

When reagents are removed from the Panther Fusion system, return them immediately to their appropriate storage temperatures.

- B. Working Panther Fusion Capture Reagent-S and Panther Fusion Enhancer Reagent-S are stable for 60 days when capped and stored at 15°C to 30°C. Do not refrigerate.
- C. Discard any unused reagents that have surpassed their on board stability.
- D. Controls are stable until the date indicated on the vials.
- E. Avoid cross-contamination during reagent handling and storage.
- F. Do not freeze reagents.

¹ On board stability starts at the time the reagent is placed on the Panther Fusion system for the Panther Fusion AdV/hMPV/RV assay cartridge, FCR-S, FER-S and IC-S. The on board stability for the Panther Fusion Reconstitution Buffer I, Panther Fusion Elution Buffer and Panther Fusion Oil Reagent starts when the reagent pack is first used.

² If removed from the Panther Fusion System, store the assay cartridge in an air-tight container with desiccant at the recommended storage temperature.

Specimen Collection and Storage

Specimens - Clinical material collected from patient placed in an appropriate transport system. For the Panther Fusion AdV/hMPV/RV assay this includes NP swab specimens in viral transport medium (VTM).

Samples - Represents a more generic term to describe any material for testing on the Panther Fusion System including specimens, specimens transferred into a Panther Fusion Specimen Lysis Tube and controls.

Note: Handle all specimens as if they contain potentially infectious agents. Use Universal Precautions.

Note: Take care to avoid cross-contamination during specimen handling steps. For example, discard used material without passing over open tubes.

A. Specimen types include NP swab specimens.

Collect NP swab specimens according to standard technique using a polyester-, rayon-, or nylon-tipped swab. Immediately place the swab specimen into 3mL of VTM.

The following types of VTM were verified for use.

- Remel MicroTest M4, M4RT, M5 or M6 formulations
- Copan Universal Transport Medium
- BD Universal Viral Transport Medium

B. Specimen processing

- 1. Prior to testing on the Panther Fusion system, transfer specimen* to a Panther Fusion Specimen Lysis Tube.
 - Transfer 500 μL of the NP swab specimens to a Panther Fusion Specimen Lysis Tube.
 - *Note: When testing frozen specimen, allow specimen to reach room temperature prior to processing.
- 2. Storing specimens before testing
 - a. After collection, specimens can be stored at 2°C to 8°C up to 96 hours before transferred to the Panther Fusion Specimen Lysis Tube. Remaining specimen volumes can be stored at ≤-70°C.
 - b. Specimen in the Panther Fusion Specimen Lysis Tube may be stored under one of the following conditions:
 - 15°C to 30°C up to 6 days or
 - 2°C to 8°C up to 3 months.

Note: It is recommended that specimens transferred to the Panther Fusion Specimen Lysis Tube are stored capped and upright in a rack.

C. Specimen on board the Panther Fusion system may be archived for additional testing at a later time.

- D. Storing samples after testing
 - 1. Samples that have been assayed should be stored upright in the rack under one of the following conditions:
 - 15°C to 30°C up to 6 days or
 - 2°C to 8°C up to 3 months.
 - 2. The samples should be covered with a new, clean plastic film or foil barrier.
 - 3. If assayed samples need to be frozen or shipped, remove the penetrable cap and place a new non-penetrable cap on the specimen tubes. If samples need to be shipped for testing at another facility, recommended temperatures must be maintained. Prior to uncapping, specimen transport tubes must be centrifuged for 5 minutes at 420 Relative Centrifugal Force (RCF) to bring all of the liquid down to the bottom of the tube. Avoid splashing and cross-contamination.

Specimen Transport

Maintain specimen storage conditions as described in the *Specimen Collection and Storage* section on page 7.

Note: Specimens must be shipped in accordance with applicable national, international, and regional transportation regulations.

Panther Fusion System

The Panther Fusion System is an integrated nucleic acid testing system that fully automates all steps necessary to perform various Panther Fusion assays from sample processing through amplification, detection, and data reduction.

Reagents and Materials Provided for Panther Fusion AdV/hMPV/RV Assay Assay Packaging

Components ¹	Part No.	Storage
Panther Fusion AdV/hMPV/RV Assay Cartridges 96 Tests Panther Fusion AdV/hMPV/RV assay cartridge, 12 tests, 8 per box	PRD-04330	2°C to 8°C
Panther Fusion Internal Control-S 960 Tests Panther Fusion Internal Control-S tube, 4 per box	PRD-04332	2°C to 8°C
Panther Fusion AdV/hMPV/RV Assay Controls Panther Fusion AdV/hMPV/RV Positive Control tube, 5 per box Panther Fusion Negative Control tube, 5 per box	PRD-04338	2°C to 8°C
Panther Fusion Extraction Reagent-S 960 Tests Panther Fusion Capture Reagent-S bottle, 240 tests, 4 per box Panther Fusion Enhancer Reagent-S bottle, 240 tests, 4 per box	PRD-04331	15°C to 30°C
Panther Fusion Elution Buffer 2400 Tests Panther Fusion Elution Buffer pack, 1200 tests, 2 per box	PRD-04334	15°C to 30°C
Panther Fusion Reconstitution Buffer I 1920 Tests Panther Fusion Reconstitution Buffer I pack, 960 tests, 2 per box	PRD-04333	15°C to 30°C
Panther Fusion Oil Reagent 1920 Tests Panther Fusion Oil Reagent pack, 960 tests, 2 per box	PRD-04335	15°C to 30°C

¹ Components can also be ordered in the following bundles: Panther Fusion Universal Fluids Kit, PRD-04430, contains 1 each Panther Fusion Oil and Panther Fusion Elution buffer. Panther Fusion Assay Fluids I-S, PRD-04431, contains 2 Panther Fusion Extraction Reagents-S, 2 Panther Fusion Internal Control-S, and 1 Panther Fusion Reconstitution Buffer I.

Individually Packaged Items

Items	Part No.
Panther Fusion Specimen Lysis Tubes, 100 per bag	PRD-04339

Materials Required and Available Separately

Note: Materials available from Hologic have catalog numbers listed, unless otherwise specified.

Material	Cat. No.
Panther System	303095
Panther Fusion Module	ASY-09600
Aptima Assay Fluids Kit (Aptima Wash Solution, Aptima Buffer for Deactivation Fluid, and Aptima Oil Reagent)	303014 (1000 tests)
Multi-tube units (MTUs)	104772-02
Panther Waste Bag Kit	902731
Panther Waste Bin Cover	504405
Or Panther System Run Kit for Real Time Assays contains MTUs, waste bags, waste bin covers, and assay fluids	PRD-03455 (5000 tests)
Or Panther System Run Kit (when running TMA assays in parallel with real time-TMA assays) contains MTUs, waste bags, waste bin covers, auto detect*, and assay fluids	303096 (5000 tests)
Panther Fusion Tube Trays, 1008 tests, 18 trays per box	PRD-04000
Liquid Handling (LiHa) Disposable Tips, 1000 μL	10612513 (Tecan)
Aptima penetrable caps (optional)	105668
Replacement non-penetrable caps (optional)	103036A
Replacement extraction reagent bottle caps	CL0040
P1000 pipettor and tips with hydrophobic plugs	-
Bleach, 5% to 7% (0.7 M to 1.0 M) sodium hypochlorite solution Note : Mix one part bleach with one part deionized water to make diluted working bleach solution 2.5% to 3.5% (0.35 M to 0.5 M) sodium hypochlorite solution.	-
Disposable powderless gloves	-

^{*}Needed only for Panther Aptima TMA assays.

Panther Fusion System Test Procedure

Note: Refer to the Panther Fusion System Operator's Manual for additional procedural information.

A. Work Area Preparation

- 1. Wipe down work surfaces with 2.5% to 3.5% (0.35 M to 0.5 M) sodium hypochlorite solution. Allow the sodium hypochlorite solution to contact surfaces for at least 1 minute and follow with a deionized (DI) water rinse. Do not allow the sodium hypochlorite solution to dry. Cover the bench surface with clean, plastic-backed absorbent laboratory bench covers.
- 2. Clean a separate work surface where samples will be prepared using the procedure described in step A.1.

B. Reagent Preparation

- 1. Remove the bottles of IC-S, FCR-S and FER-S from storage.
- 2. Open the bottles of IC-S, FCR-S and FER-S, and discard the caps. Open the TCR door on the upper bay of the Panther Fusion system.
- 3. Place the IC-S, FCR-S and FER-S bottles in the appropriate positions on the TCR carousel.
- 4. Close the TCR door.

Note: The Panther Fusion system adds the IC-S to the FCR-S. After the IC-S is added to the FCR-S, it is referred to as wFCR-S (working FCR-S). If the FCR-S and FER-S are removed from the system, use new caps and immediately store according to the proper storage conditions.

C. Specimen Handling

Note: Prepare specimens per the Specimen Processing instructions in the Specimen Collection and Storage section before loading specimens onto the Panther Fusion system.

- 1. Do not vortex samples.
- Inspect sample tubes before loading into the rack. If a sample tube contains bubbles or has a lower volume than is typically observed, gently tap the bottom of the tube to bring contents to the bottom.

Note: To avoid a processing error, ensure adequate specimen volume is added to the Panther Fusion Specimen Lysis Tube. When 500 μ L of NP swab specimen is added to the Panther Fusion Specimen Lysis Tube, there is sufficient volume to perform 3 nucleic acid extractions.

D. System Preparation

For instructions on setting up the Panther Fusion system including loading samples, assay cartridges and universal fluids, refer to the *Panther Fusion System Operator's Manual*.

Procedural Notes

A. Controls

- 1. The Panther Fusion AdV/hMPV/RV Positive Control and Panther Fusion Negative Control can be loaded in any rack position, in any Sample Bay lane on the Panther Fusion system.
- 2. Once the control tubes are pipetted and are processed for the Panther Fusion AdV/hMPV/RV assay, they are active for up to 30 days (control frequency configured by an administrator) unless control results are invalid or a new assay cartridge lot is loaded.
- 3. Each control tube can be tested once.
- 4. Patient specimen pipetting begins when one of the following two conditions is met:
 - a. Valid results for the controls are registered on the system.
 - b. A pair of controls is currently in process on the system.

Quality Control

A run or specimen result may be invalidated by the Panther Fusion System if problems occur while performing the assay. Specimens with invalid results must be retested.

Negative and Positive Controls

To generate valid results, a set of assay controls must be tested. One replicate of the negative assay control and positive assay control must be tested each time a new lot of assay cartridges is loaded on the Panther Fusion system or when the current set of valid controls for an active cartridge lot have expired.

The Panther Fusion system is configured to require assay controls run at an administrator-specified interval of up to 30 days. Software on the Panther Fusion system alerts the operator when assay controls are required and does not start new tests until the assay controls are loaded and have started processing.

During processing, criteria for acceptance of the assay controls are automatically verified by the Panther Fusion system. To generate valid results, the assay controls must pass a series of validity checks performed by the Panther Fusion system.

If the assay controls pass all validity checks, they are considered valid for the administrator-specified time interval. When the time interval has passed, the assay controls are expired by the Panther Fusion system and requires a new set of assay controls be tested prior to starting any new samples.

If any one of the assay controls fails the validity checks, the Panther Fusion system automatically invalidates the affected samples and requires a new set of assay controls be tested prior to starting any new samples.

Internal Control

An internal control is added to each sample during the extraction process. During processing, the internal control acceptance criteria is automatically verified by the Panther Fusion system software. Detection of the internal control is not required for samples that are positive for AdV, hMPV and/or RV. The internal control must be detected in all samples that are negative for AdV, hMPV and RV targets; samples that fail to meet that criteria will be reported as Invalid. Each sample with an Invalid result must be retested.

The Panther Fusion system is designed to accurately verify processes when procedures are performed following the instructions provided in this package insert and the *Panther Fusion System Operator's Manual*.

Interpretation of Results

The Panther Fusion system automatically determines the test results for samples and controls. Results for AdV, hMPV and RV detection are reported separately. A test result may be negative, positive, or invalid.

Table 1 shows the possible results reported in a valid run with result interpretations.

Table 1: Result Interpretation

AdV Result	hMPV Result	RV Result	IC Result	Interpretation
Neg	Neg	Neg	Valid	AdV, hMPV and RV not detected.
POS	Neg	Neg	Valid	AdV detected. hMPV and RV not detected.
Neg	POS	Neg	Valid	hMPV detected. AdV and RV not detected.
Neg	Neg	POS	Valid	RV detected. AdV and hMPV not detected.
POS	POS	Neg	Valid	AdV and hMPV detected. RV not detected.
Neg	POS	POS	Valid	hMPV and RV detected. AdV not detected.
POS	Neg	POS	Valid	AdV and RV detected. hMPV not detected.
POS	POS	POS	Valid	AdV, hMPV and RV detected. Triple infections are rare. Retest to confirm result.
Invalid	Invalid	Invalid	Invalid	Invalid. There was an error in the generation of the result; retest sample.

Note: POS result will be accompanied by cycle threshold (Ct) values.

Limitations

- A. Use of this assay is limited to personnel who are trained in the procedure. Failure to follow these instructions may result in erroneous results.
- B. Reliable results are dependent on adequate specimen collection, transport, storage, and processing.
- C. Avoid contamination by adhering to good laboratory practices and to the procedures specified in this package insert.
- D. Negative results do not preclude adenovirus, human metapneumovirus or rhinovirus infections and should not be used as the sole basis for treatment or other management decisions.
- E. This test does not differentiate Adenovirus subtypes (i.e. 1-58), human Metapneumovirus subtypes (i.e., A1, A2, B1, B2) or Rhinovirus species (i.e., Rhinovirus A, Rhinovirus B or Rhinovirus C); additional testing is required to differentiate any specific Adenovirus subtypes, human Metapneumovirus subtypes or specific Rhinovirus species in consultation with local public health departments.
- F. A positive result indicates the detection of nucleic acid from the relevant virus. Nucleic acid may persist even after the virus is no longer viable.

Panther Fusion System Assay Performance

Clinical Performance

Retrospectively collected NP swabs from patients in the US with reference test results were used for evaluation. The results are shown in tables 2, 3, and 4.

For NP swab specimens, 500 microliter (μ L) was diluted into a Panther Fusion Specimen Lysis Tube containing 780 μ L of Specimen Transport Media (STM) and a single replicate was tested with the Panther Fusion AdV/hMPV/RV assay. The result was compared to an FDA cleared nucleic acid test (NAT) result. The sensitivity and specificity for the detection of AdV, hMPV, and RV nucleic acid was determined.

A total of 546 NP swab specimens were tested with the Panther Fusion AdV/hMPV/RV assay and with Luminex xTAG® Respiratory Viral Panel or Luminex xTAG® Respiratory Viral Panel FAST v2 or GenMark Dx eSensor Respiratory Viral Panel. Sensitivity and specificity for detection of AdV, hMPV, and RV are shown for NP swab specimens.

Table 2: AdV Results

Specimen Type		AdV+	AdV-		- Sensitivity	Specificity	Overall	
Specimen Type	N	Fusion AdV	Fusion AdV	Fusion AdV	Fusion AdV	95% CI	95% CI	Agreement 95% CI
		+	-	+	-			
Nasopharyngeal	546	175	3*	11**	357	98.3%	97.0%	97.4%
Swab	540	173	75 5	11	331	95.2 - 99.4%	94.7 - 98.3%	95.7 - 98.5%

^{*} Two out of three discordant specimens were tested with FDA cleared assay. AdV was not detected in both specimens. Untested discordant specimens had insufficient volumes.

Table 3: hMPV Results

Specimen Type		hM	PV+	hM	PV-	0 10110	0	Overall
Specimen Type	N	Fusion hMPV	Fusion hMPV	Fusion hMPV	Fusion hMPV	Sensitivity95% CI	Specificity 95% CI	Agreement 95% CI
		+	-	+	-			
Nasopharyngeal	546	104	0	24*	418	100.0%	94.6%	95.6%
Swab	340	104	U	24	410	96.4 - 100.0%	92.0- 96.3%	93.5 - 97.0%

^{*} Nineteen out of 24 discordant specimens were tested with in-house developed and validated RT-PCR assay. hMPV was detected in four specimens. Untested discordant specimens had insufficient volumes.

^{**} Six out of eleven discordant specimens were tested with FDA cleared assay. AdV was detected in five specimens. Untested discordant specimens had insufficient volumes.

Table 4: RV Results

Specimen Type		RV+ RV-			Sensitivity	Specificity	Overall	
Specimen Type	N	Fusion RV	Fusion RV	Fusion RV	Fusion RV	95% CI	95% CI	Agreement 95% CI
		+	-	+	-	_		
Nasopharyngeal Swab	546	255	28*	12**	251	90.1%	95.4%	92.7%
Nasopharyngear Swab	340	200	20	12	231	86.1 - 93.1%	92.2 - 97.4%	90.2 - 94.6%

^{*} Twenty three out of 28 discordant specimens were tested with an in-house developed and validated bidirectional sequencing assay. RV was not detected in 16 out of 23 tested. Untested discordant specimens had insufficient volumes.

Analytical Sensitivity

The analytical sensitivity (limit of detection or LoD) of the Panther Fusion AdV/hMPV/RV assay for the NP swab specimen type was determined by testing pooled AdV/hMPV/RV negative clinical specimens spiked with the following virus cultures at various concentrations: Adenovirus (1, 3, 4, 9, 12, 40), hMPV (A1, A2, B1, B2) and RV (A-18 and B-26). At least twelve replicates were tested with each of the three reagent lots for a combined total of 36 replicates. Target specific LoD concentrations were verified by testing an additional 20 replicates with one reagent lot. Analytical sensitivity (LoD) is defined as the lowest concentration at which ≥95% of all replicates tested positive, as summarized in the table below.

Table 5: NP Swab Sensitivity

Viral Strain	LoD Concentration
Adenovirus 1 (Species C)	1x10º TCID50/mL
Adenovirus 3 (Species B)	1x10º TCID50/mL
Adenovirus 4 (Species E)	1x10-2 TCID50/mL
Adenovirus 9 (Species D)	1x10 ^{-0.5} TCID ₅₀ /mL
Adenovirus 12 (Species A)	1x10 ^{-0.5} TCID ₅₀ /mL
Adenovirus 40 (Species F)	1x10 ^{-1.5} TCID ₅₀ /mL
hMPV A1-16	1x10 ² TCID ₅₀ /mL
hMPV A2-20	1x10 ¹ TCID ₅₀ /mL
hMPV B1-3	1x10 ^{0.5} TCID ₅₀ /mL
hMPV B2-8	1x10 ⁰ TCID ₅₀ /mL
Rhinovirus A-18	1x10 ^{-0.5} TCID ₅₀ /mL
Rhinovirus B-26	1x10º TCID50/mL

^{**} All 12 discordant specimens were tested with an in-house developed and validated bidirectional sequencing assay. RV was detected in nine specimens.

Reactivity

The reactivity of the Panther Fusion AdV/hMPV/RV assay was evaluated against multiple strains of AdV, hMPV, and RV. Simulated reactivity evaluation was performed *in silico* for the types that are not available for testing. Reactivity was predicted for the AdV type 52-58 and RV type C.

Table 6: Reactivity Results

Target	Description	Concentration	AdV	hMPV	RV
Adenovirus	AdV 1	1x103 TCID50/mL	+	-	-
	AdV 2	1x10³ TCID₅₀/mL	+	-	-
	AdV 3	1x10³ TCID₅₀/mL	+	-	-
	AdV 4	1x103 TCID50/mL	+	-	-
	AdV 5	1x10³ TCID50/mL	+	-	-
	AdV 6	1x10³ TCID₅₀/mL	+	-	-
	AdV 7	1x103 TCID50/mL	+	-	-
	AdV 8	1x10³ TCID₅₀/mL	+	-	-
	AdV 9	1x10³ TCID₅₀/mL	+	-	-
	AdV 10	1x103 TCID50/mL	+	-	-
	AdV 11	1x10³ TCID₅₀/mL	+	-	-
	AdV 12	1x10³ TCID₅₀/mL	+	-	-
	AdV 13	1x103 TCID50/mL	+	-	-
	AdV 14	1x10³ TCID₅₀/mL	+	-	-
	AdV 15	1x10³ TCID₅₀/mL	+	-	-
	AdV 16	1x10³ TCID₅₀/mL	+	-	-
	AdV 17	1x10³ TCID₅₀/mL	+	-	-
	AdV 19	1x10³ TCID₅₀/mL	+	-	-
	AdV 20	1x10³ TCID₅₀/mL	+	-	-
	AdV 21	1x10³ TCID₅₀/mL	+	-	-
	AdV 22	1x10³ TCID₅₀/mL	+	-	-
	AdV 23	1x10³ TCID50/mL	+	-	-
	AdV 24	1x10³ TCID₅₀/mL	+	-	-
	AdV 25	1x10³ TCID₅₀/mL	+	-	-
	AdV 26	1x10³ TCID50/mL	+	-	-
	AdV 27	1x10³ TCID₅₀/mL	+	-	-
	AdV 28	1x10³ TCID₅₀/mL	+	-	-
	AdV 29	1x10³ TCID50/mL	+	-	-
	AdV 30	1x10³ TCID₅₀/mL	+	-	-
	AdV 31	1x10³ TCID₅₀/mL	+	-	-
	AdV 32	1x103 TCID50/mL	+	-	-
	AdV 33	1x10³ TCID₅₀/mL	+	-	-
	AdV 34	1x10³ TCID₅₀/mL	+	-	-
	AdV 35	1x10³ TCID₅₀/mL	+	_	

Table 6: Reactivity Results (continued)

Target	Description	Concentration	AdV	hMPV	RV
Adenovirus	AdV 36	1x103 TCID50/mL	+	-	-
	AdV 37	1x103 TCID50/mL	+	-	-
	AdV 38	1x103 TCID50/mL	+	-	-
	AdV 39	1x103 TCID50/mL	+	-	-
	AdV 40	1x103 TCID50/mL	+	-	-
	AdV 41	1x103 TCID50/mL	+		-
	AdV 42	1x103 TCID50/mL	+	-	-
	AdV 43	1x103 TCID50/mL	+	-	-
	AdV 44	1x103 TCID50/mL	+	-	-
	AdV 45	1x103 TCID50/mL) ³ TCID ₅₀ /mL + -		-
	AdV 46	1x103 TCID50/mL			-
	AdV 47	1x103 TCID50/mL	+	-	-
	AdV 48	1x103 TCID50/mL	+	-	-
	AdV 49	1x103 TCID ₅₀ /mL	+	-	-
	AdV 50	1x103 TCID50/mL	+	-	-
	AdV 51	1x103 TCID50/mL	+	-	-
human	hMPV A1-16	1x10 ³ TCID ₅₀ /mL	-	+	-
Metapneumovirus	hMPV A1-9	1x103 TCID50/mL	-	+	-
	hMPV A2-20	1x103 TCID50/mL	-	+	-
	hMPV A2-27	1x10 ³ TCID ₅₀ /mL	-	+	-
	hMPV B1-3	1x103 TCID50/mL	-	+	-
	hMPV B1-5	1x103 TCID50/mL	-	+	-
	hMPV B2-18	1x103 TCID ₅₀ /mL	-	+	-
	hMPV B2-4	1x103 TCID50/mL	-	+	-
	hMPV B2-8	1x103 TCID50/mL	-	+	-
Rhinovirus*	RV A1	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A16	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A18	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A32	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A33	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A39	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A40	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A44	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A51	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A59	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A61	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A65	1x10 ² TCID ₅₀ /mL	-	-	+

Table 6: Reactivity Results (continued)

Target	Description	Concentration	AdV	hMPV	RV
Rhinovirus*	RV A76	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A78	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A89	1x10 ² TCID ₅₀ /mL	-	-	+
	RV A100	1x10 ² TCID ₅₀ /mL	-	-	+
	RV B26	26 1x10² TCID₅₀/mL		-	+
	RV B52	RV B52 1x10 ² TCID ₅₀ /mL -		-	+
	RV B69 1x10 ² TCID ₅₀ /ml		-	-	+
	RV B70	1x10 ² TCID ₅₀ /mL	-	-	+
	RV B79	1x10² TCID₅₀/mL	-	-	+
	RV B86	1x10² TCID₅₀/mL	-	-	+

^{*} Simulated reactivity evaluation performed in-silico predicted reactivity with multiple Rhinovirus C strains.

Analytical Specificity

The analytical specificity of the Panther Fusion AdV/hMPV/RV assay was evaluated by testing a panel of 64 organisms, consisting of 30 viral, 32 bacterial, and 2 yeast strains representing common respiratory pathogens or flora commonly present in nasopharynx. Bacteria and yeast were tested at concentrations of 10⁵ to10⁸ CFU/mL or IFU/mL, except where noted. Viruses were tested at concentrations of 10³ to 10⁷ TCID₅₀/mL.

Analytical specificity of the Panther Fusion AdV/hMPV/RV assay was 100% for AdV, hMPV, and RV.

Table 7: Specificity Results

Organism	Concentration	AdV	hMPV	RV
Acinetobacter baumannii 307-0294	1x10 ⁷ CFU/mL	-	-	-
Bordetella bronchiseptica	1x10 ⁷ CFU/ml	-	-	-
Bordetella parapertussis	1x10 ⁷ CFU/ml	-	-	-
Bordetella pertussis	1x107 CFU/mL	-	-	-
Burkholderia cepacia Z066	1x10 ⁶ CFU/mL	-	-	-
Candida albicans	1x10 ⁷ CFU/mL	-	-	-
Candida glabrata	1x10 ⁶ CFU/mL	-	-	-
Chlamydia pneumoniae	1x10⁵ CFU/mL	-	-	-
Chlamydia trachomatis	1x10⁴ CFU/mL	-	-	-
CMV Strain AD 169	1x10⁴ TCID₅₀/mL	-	-	-
Coronavirus 229E	1x10⁵ TCID₅₀/mL	-	-	-
Coronavirus OC43	1x10⁵ TCID₅₀/mL	-	-	-
Corynebacterium diphtheria	1x107 CFU50/mL	-	-	-
Coxsackie B3	1x106 TCID₅₀/mL	-	-	-
Coxsackie B4	1x10 ⁴ TCID ₅₀ /mL	-	-	-

Table 7: Specificity Results (continued)

Organism	Concentration	AdV	hMPV	RV	
Coxsackie B5/10/2006	1x10⁵ TCID₅₀/mL	-	-	-	
Coxsackievirus A10	1x10 ⁴ TCID ₅₀ /mL	-	-	-	
Coxsackievirus A21	1x10 ⁴ TCID ₅₀ /mL	-	-	-	
E. coli	1x10 ⁷ CFU/mL	-	-	-	
EBV	1x106 TCID₅o/mL	-	-	-	
Echovirus 11	1x10 ⁶ TCID₅₀/mL	-	-	-	
Echovirus 2	1x10 ⁶ TCID ₅₀ /mL	-	-	-	
Echovirus 3	1x10⁴ TCID₅₀/mL	-	-	_	
Echovirus 6	1x106 TCID50/mL	_	-	_	
Enterovirus 68	1x10⁵ TCID₅₀/mL	_	-	_	
Enterovirus 70	1x10 ⁴ TCID ₅₀ /mL	_	_	_	
Haemophilus Influenzae	1x10 ⁷ TCID ₅₀ /mL	_	_	_	
HPIV-1	1x10 ⁴ TCID ₅₀ /mL	_			
HPIV-2	1x10 TCID50/mL		-	-	
		-	-	-	
HPIV-3	1x10 ⁵ TCID ₅₀ /mL	-	-	-	
HPIV-4a	1x10⁴ TCID₅₀/mL	-	-	-	
HSV-1 Macinytre Strain	1x10⁵ TCID₅₀/mL	-	-	-	
HSV-2 Type 2G Strain	1x10⁵ TCID₅₀/mL	-	-	-	
Influenza A (H1N1)	1x10 ³ TCID ₅₀ /mL	-	-	-	
Influenza A (H3N2)	1x103 TCID50/mL	-	-	-	
Influenza B	1x10 ³ TCID ₅₀ /mL	-	-	-	
Klebsiella pneumonia	1x10 ⁷ CFU/mL	-	-	-	
Lactobacillus acidophilus Z048	1x10 ⁶ CFU/mL	-	-	-	
Lactobacillus plantarum	1x10 ⁶ CFU/mL	-	-	-	
Legionella pneumophila	1x10 ⁷ CFU/mL	-	-	-	
Measles/7/2000	1x10⁴ TCID₅₀/mL	-	-	-	
Moraxella catarrhalis	1x10 ⁷ CFU/mL	-	-	-	
Mumps virus	1x10⁵ CFU/mL	-	-	-	
Mycobacterium intracellulare	5x10 ¹⁰ rRNA copies/mL	-	-	-	
Mycobacterium tuberculosis	5x10° rRNA copies/mL	-	-	-	
Mycoplasma pneumoniae	1x10 ⁶ CFU/mL	-	-	-	
Neisseria gonorrhea	1x10 ⁷ CFU/mL	-	-	-	
Neisseria meningitides	1x10 ⁷ CFU/mL	-	-	-	
Neisseria mucosa	1x10 ⁷ CFU/mL	-	-	-	
Polio virus 1	1x106 TCID50/mL	_	_	_	

Table 7: Specificity Results (continued)

Organism	Concentration	AdV	hMPV	RV
Proteus mirabilis	1x10 ⁷ CFU/mL	-	-	-
Proteus vulgaris	1x10 ⁷ CFU/mL	-	-	-
Pseudomonas aeruginosa	1x10 ⁷ CFU/mL	-	-	-
RSV A	1x10⁵ TCID₅₀/mL	-	-	-
RSV B	1x10⁵ TCID₅₀/mL	-	-	-
Serratia marcescens Z053	1x10 ⁷ CFU/mL	-	-	-
Staphlycoccus aureus	1x10 ⁷ CFU/mL	-	-	-
Staphlycoccus epidermidis	1x10 ⁷ CFU/mL	-	-	-
Streptococcus agalactiae	1x10 ⁷ CFU/mL	-	-	-
Streptococcus pneumoniae	1x10 ⁷ CFU/mL	-	-	-
Streptococcus pyogenes	1x10 ⁷ CFU/mL	-	-	-
Streptococcus salivarius	1x10 ⁷ CFU/mL	-	-	-
Tatlockia micdadei (Legionella micdadei)	1x10 ⁶ CFU/mL	-	-	-
Varicella Zoster Virus	1x10⁴ TCID₅₀/mL	-	-	-

Competitive Interference

Competitive Interference of the Panther Fusion AdV/hMPV/RV assay was evaluated using a simulated clinical matrix with pairs of target viruses at two different concentrations. One of the concentrations was near the Limit of Detection (3X LoD) while the other concentration was high (1000X LoD). The presence of two viruses at varying concentrations in a single sample had no effect on the analytical sensitivity (100% detection for both targets) at the concentration noted in the table below.

Table 8: Competitive Interference

Condition	Tar	get 1	Tar	AdV	hMPV	RV	
	Description	Concentration	Description	Concentration	Result	Result	Result
1	AdV	3X LoD	hMPV	1000X LoD	+	+	-
2	AdV	3X LoD	RV 1000X LoD		+	-	+
3	hMPV	3X LoD	AdV	1000X LoD	+	+	-
4	hMPV	3X LoD	RV 1000X LoD		-	+	+
5	RV	3X LoD	AdV	1000X LoD	+	-	+
6	RV	3X LoD	hMPV	1000X LoD	ı	+	+

Interference

Mucin, whole blood and other potentially interfering substances (medications and over-the-counter or OTC products) that may be present in the samples were evaluated in the Panther Fusion AdV/hMPV/RV assay. Clinically relevant amount of the potentially interfering substances were added to simulated clinical matrix and tested unspiked or spiked with cultured AdV, hMPV

and RV at their respective 3X LoD concentrations. The substances consisted of nasal sprays (liquid and powder), ingestible pills, lozenges, injectable and endogenous substances, as shown in Table 9.

All of the substances tested were found to have no impact on the performance of the Panther Fusion AdV/hMPV/RV assay.

Table 9: Potentially Interfering Substances

Туре	Substance Name	Active Ingredient(s)	Concentration
Endogenous	Mucin	Purified mucin protein	60 μg/mL
Endogenous	Human blood	Blood	2% v/v
	Neo-Synephrine®	Phenylephrine	15% v/v
Nasal sprays or drops	Anefrin	Oxymetazoline	15% v/v
Nasai spiays of diops	Saline	Sodium chloride	15% v/v
	Ventolin® HFA	Albuterol	15% v/v
	QVAR®, Beconase AQ	Beclomethasone	5% v/v
	Dexacort	Dexamethasone	5% v/v
	AEROSPAN®	Flunisolide	5% v/v
Nasal corticosteroids	Nasacort	Triamcinolone	5% v/v
	Rhinocort	Budesonide	5% v/v
	Nasonex	Mometasone	5% v/v
	Flonase	Fluticasone	5% v/v
Nasal gel	Zicam® (Allergy Relief)	Luffa opperculata, Galphimia, Glauca, Histaminum hydrochloricum, Sulfur	5% v/v
Throat lozenges	Chloraseptic Throat Lozenges	Benzocaine Menthol	0.63 mg/mL
	Relenza®	Zanamivir	3.3 mg/mL
Anti-viral drugs	TamiFlu	Oseltamivir	25 mg/mL
	Rebitol	Ribavirin	20 mg/mL
Antibiotic, nasal ointment	Bactroban cream	Mupirocin	10 mg/mL
Antibiotic, systemic	Tobramycin	Tobramycin	4.0 μg/mL

Carry-Over/Contamination

The carry-over/cross-contamination study was performed with negative samples alternately placed between high positive samples and tested. High positive samples were prepared by spiking (over 10,000X LoD). Nine separate runs with negative samples and positive samples placed in a checkerboard pattern were tested over three different instruments for a combined total of 449 positive and 450 negative samples. The carry-over rate was 0.2%.

Assay Precision

Panther Fusion AdV/hMPV/RV assay precision was evaluated with a 7-member panel. The panel was tested by three operators on two separate runs per day, using three reagent lots on three Panther Fusion systems over 45 days.

The panel members are described in Table 10, along with a summary of the agreement with expected results for each targets. Table 11 presents the mean and variability analysis between instruments, between reagent lots, between operators, between days, between runs and within runs, and overall (total) for Ct.

Table 10: Panel Description and % Agreement

Target	Panel Member	% Positive	% Total Agreement (95% CI)
	AdV	100.0%	100.0%
	3x LoD	(162/162)	(97.7 - 100%)
AdV	AdV	100.0%	100.0%
	1x LoD	(162/162)	(97.7 - 100%)
Auv	AdV	10.6%	89.4%
	0.01x LoD	(17/161)	(83.7 - 93.3%)
	Negative	0.6% (1/162)	99.4% (96.6 - 99.9%)
	hMPV	100.0%	100.0%
	3x LoD	(160/160)	(97.7 - 100%)
hMPV	hMPV	100.0%	100.0%
	1x LoD	(161/161)	(97.7 - 100%)
HIMPV	hMPV	17.9%	82.1%
	0.01x LoD	(29/162)	(75.5 - 87.2%)
	Negative	0.0% (0/162)	100.0% (97.7% - 100.0%)
	RV	100.0%	100.0%
	3x LoD	(161/161)	(97.7 - 100%)
RV	RV	100.0%	100.0%
	1x LoD	(162/162)	(97.7 - 100%)
I KV	RV	1.9%	98.1%
	0.01x LoD	(3/160)	(94.6 - 99.4%)
	Negative	0.6% (1/162)	99.4% (96.6 - 99.9%)

Table 11: Signal Variability

Target	Panel Member	Mean Ct		ween ument		ween ent Lots		ween erator		ween ays	Betwee	en Runs	Withi	n Runs	T	otal
			SD	CV (%)	SD	CV (%)	SD	CV (%)	SD	CV (%)	SD	CV (%)	SD	CV (%)	SD	CV (%)
	AdV 3x LoD	33.5	0.1	0.4	0.0	0.1	0.0	0.0	0.1	0.3	0.2	0.7	0.4	1.2	0.5	1.5
AdV	AdV 1x LoD	35.2	0.2	0.6	0.0	0.0	0.0	0.2	0.1	0.3	0.3	0.8	0.5	1.5	0.6	1.9
	AdV 0.01x LoD	40.4	0.3	0.8	0.0	0.0	0.0	0.0	0.0	0.0	0.9	2.4	0.7	1.9	1.3	3.2
	hMPV 3x LoD	33.5	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.2	0.8	0.8	2.4	0.8	2.5
hMPV	hMPV 1x LoD	35.1	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.7	2.0	0.7	2.0
	hMPV 0.01x LoD	40.3	0.0	0.0	0.0	0.0	0.0	0.0	0.3	0.7	0.5	1.4	1.2	3.1	1.4	3.5
	RV 3x LoD	32.5	0.1	0.5	0.1	0.3	0.0	0.1	0.0	0.0	0.3	1.0	0.6	2.0	0.7	2.4
RV	RV 1x LoD	33.8	0.1	0.5	0.1	0.5	0.0	0.0	0.1	0.4	0.0	0.0	0.8	2.6	0.9	2.8
	RV 0.01x LoD	40.6	1.9	4.7	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.6	1.6	2.0	5.0
IC	Negative	30.7	0.0	0.2	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.6	0.5	1.7	0.5	1.8

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