

CLIA Complexity: Waived

INTENDED USE

The Clearview RSV test is intended for the rapid, qualitative detection of respiratory syncytial virus fusion protein directly from nasopharyngeal swab, and nasal aspirate specimens in children less than 6 years and adults over the age of 60. The test is intended for use as an aid in the rapid laboratory diagnosis of acute respiratory syncytial virus infection in patients with symptoms consistent with RSV infection. It is recommended that negative test results be confirmed by cell culture or DFA.

SUMMARY AND EXPLANATION

Respiratory syncytial virus is a highly contagious, acute, viral infection of the respiratory tract. The causative agent is a single stranded (negative strand) RNA virus of the paramyxoviridae family. RSV is now recognized as the leading cause of hospitalization of children during the first year of life. It is also the major viral cause of nosocomial illness in children already hospitalized for other reasons. Half of all infants become infected during their first year of life. Virtually all have been infected by their second year.¹ Infection involving the lower respiratory tract carries an associated mortality rate of 0.5%, especially in premature infants or infants and children with underlying lung disease.

RSV antigens may be detected in clinical specimens by im-

munoassay. The Clearview RSV Test is a gold-labeled lateral flow immunoassay using monoclonal antibodies directed against RSV fusion (F) protein antigens.

PRINCIPLE OF THE TEST

The Clearview RSV is a lateral flow immunogold assay for the direct visual detection of RSV protein F in clinical samples. The basis for protein F detection is in the use of a red - colored gold labeled mouse monoclonal anti-RSV protein F antibody that after addition of extracted sample travels laterally along the strip test device membrane. This lateral flow carries the mixture of sample and gold labeled anti-RSV protein F through a membrane adsorbed monoclonal anti-RSV protein F Test Line (T) and then through a membrane adsorbed goat anti-mouse immunoglobulin Control Line (C). When RSV protein F is present in clinical samples, the fluid phase mouse anti-RSV protein F binds this antigen and this formation of antigen - antibody complex is then in turn bound at the Test Line (T). The unbound or excess mouse anti-RSV protein F passes through the Test Line (T) and is bound at the Control Line (C) by goat anti- mouse immunoglobulin.

Therefore, in the presence of RSV protein F antigen, 2 red lines become visible: one at the Test Line (T) and a second at the Control Line (C). But when RSV antigen is absent only one red line appears at the Control Line (C).

REAGENTS AND MATERIALS SUPPLIED

The amounts of material or reagents included depends on kit size (20 or 40 determinations) as follows:

- Individually packaged Clearview RSV test devices - 20 for 20 test kits or 40 for 40 test kits: A membrane coated with mouse monoclonal antibody specific for RSV antigen (fusion protein, IgG, 150,000 Daltons) and with control line antibody is combined with gold labeled mouse anti-RSV antibody in a lateral flow sandwich test device.
- R1 (Extraction Reagent Solution) - 2 mL for 20 tests or 4 mL for 40 tests: surfactant in 0.5 molar MOPS buffer pH 8.0
- Extraction Tubes - 20 for 20 tests and 40 for 40 tests: disposable test vials for mixing 160 uL of patient sample with 4 drops of extraction reagent.
- Disposable Pipettes - 40 for 20 tests and 80 for 40 tests: calibrated transfer pipettes used to transfer a specified amount (160 uL) of patient sample to the extraction tube and to transfer the sample-extraction reagent mixture to the test device. **Use only the pipettes provided by IBC or a calibrated pipette capable of delivering 160uL and 240 uL sample volumes.**

- PC (Positive Control Reagent) - 1.0 mL for 20 tests or 1.5 mL for 40 tests: inactivated Long Strain of RSV at 3.8×10^5 pfu/mL in Extraction Reagent
- NC (Negative Control Reagent) - 1.0 mL for 20 tests or 1.5 mL for 40 tests: – Extraction Reagent Solution
- Package insert (1)

MATERIALS NOT SUPPLIED

- Specimen containers
- Timer or watch
- Sterile Normal Saline
- Aspiration Bulb
- Vortexer or other mixing device
- Sterile Nasopharyngeal swabs (see SAMPLE TRANSPORT AND STORAGE section)

WARNINGS AND PRECAUTIONS

- The Clearview RSV Test is for *in vitro* diagnostic use.
- The positive control (PC) is made with Clorox inactivated RSV and should be handled as though it could transmit disease.
- Do not use the kit contents beyond the expiration date printed on the outside of the box and on the individual components

- Use appropriate precautions against microbial hazards in the collection, handling, storage and disposal of patient samples and used kit contents.² Discard used material in a proper biohazard or sharps container. Patient samples should be handled as though they could transmit disease.
- The test device must remain sealed in the protective foil pouch until use.
- The R1, PC and NC contain a detergent solution. If the solution contacts the skin or eye, flush with copious amounts of water.
- To obtain accurate results, you must follow the test procedure in the package insert
- To obtain accurate results, use appropriate nasopharyngeal swabs and recommended transport media.
- All transfer pipettes and test vials are single use items - do not use more than once.

KIT STORAGE AND STABILITY

Store the kit refrigerated or at room temperature, (2-30°C), out of direct sunlight. Kit contents are stable until the expiration date printed on the package and kit contents. Do not freeze. Remove the Test Device from the foil pouch just prior to use.

SPECIMEN COLLECTION

Nasopharyngeal Swab Sample:

To collect a nasopharyngeal swab sample, insert the sterile swab into the nostril that presents the most secretion under visual inspection. Using gentle rotation, push the swab until resistance is met at the level of the turbinates (less than one inch into the nostril). Rotate the swab a few times against the nasal wall. Nasopharyngeal specimens should be placed in a maximum of 3 mL normal saline or transport medium prior to processing. It is recommended that the viral transport medium or saline should be limited to no more than 3 mL total volume. For nasal swabs, Dacron™ polyester or rayon tipped swabs with an aluminum wire are recommended. See SAMPLE TRANSPORT AND STORAGE section for appropriate swabs.

Nasal Aspirate Procedure For Older Children and Adults:

Fill aspiration bulb or bulb syringe with 2.0 –2.5 mL of sterile normal saline. Insert the tip of the bulb into the nostril until the nostril seals around the bulb and instill the saline into one nostril while the head is tilted back. Release the pressure on the bulb to aspirate the specimen back into the bulb. Transfer the specimen to a clean dry specimen container. Repeat for the other nostril and collect the fluid into the same specimen container.

For Younger Children:

The child should sit on the parent's lap facing forward, with the child's back against the parent's chest. The parent should wrap one arm around the child in a manner that will restrain the child's body and arms.

Fill aspiration bulb or bulb syringe with 1.5 –2.0 mL of sterile normal saline (depending on the size of the child). Insert the tip of the bulb into the nostril until the nostril seals around the bulb and instill the saline into one nostril while the head is tilted back. Release the pressure on the bulb to aspirate the specimen back into the bulb. Transfer the specimen to a clean dry specimen container. Repeat the process for the other nostril and collect the fluid into the same specimen container.

Nasal aspirate volumes of 3 to 5 mL are recommended. Excessive sample volumes may result in decreased test performance. If viral transport medium is added to the specimen, limit the total volume to no more than 5 mL. Process specimen as described in "Test Procedures".

SAMPLE TRANSPORT AND STORAGE

Samples should be tested as soon as possible after collection. Samples may be stored refrigerated (2-8°C) in a clean, dry, closed container for up to 48 hours or frozen at –70°C for up to one week prior to processing. **Do not centrifuge specimen prior to use with the Clearview RSV Test Device, as removal of cellular material will adversely affect the sensitivity of the test.** Because RSV is a very thermolabile organism, extreme caution should be used in collection and transport with regard to time, temperature and use of recommended transport medium. The following transport media have been tested and found to be compatible with the Clearview RSV Test:

Saline, Sterile Normal
Phosphate Buffered Saline
PBS plus 0.5% gelatin
PBS plus 0.5% Bovine Serum Albumin (BSA)
Viral Culturette™ (ideal for testing a negative result)
Veal Infusion Broth (VIB)
VIB plus 0.5% BSA
Earle's Minimal Essential Medium (EMEM)
EMEM with Lactalbumin Hydrolysate
Trypticase™ Soy Broth plus 0.5% gelatin
M5 media
Microgent™ /Swab Combo in M4 media by Micro Test (ideal for testing a negative result) This swab and transport media kit was utilized in the clinical performance studies for Clearview RSV.

QUALITY CONTROL

Built-in Procedural Control Features

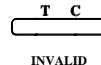
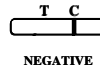
The Clearview RSV Test contains built-in procedural control features. The manufacturer's recommendations for daily control is to document these built-in procedural control features for each sample tested. The one or two line result format provides a simple interpretation for negative or positive results. The appearance of the one pink to red procedural Control Line (C) provides several forms of procedural control by demonstrating sufficient capillary flow has occurred and the functional integrity of the Test Device was maintained. **If the pink to red procedural Control Line (C) does not develop at 15 minutes, the test result is considered invalid.** Should this occur, review the test procedure and repeat the test with a new Test Device.

External Quality Control

It is recommended that external controls be used to demonstrate that the reagents and assay procedure performed properly.

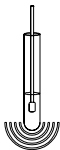
External Positive and Negative Control Reagents (PC and NC) are supplied in the Test Kits and should be tested by adding six (6) drops of the PC or NC to the Sample Window (S) on the test device and allow the test to develop as described in the TEST PROCEDURES section. **The PC and NC bottles must be swirled or gently inverted twice before being used.** These controls must be tested with each new lot or shipment of materials (20 and 40 test kits) and with each new operator. Additional controls may be tested according to guidelines or requirements of local, state, and/or federal regulations or accrediting organizations. It is recommended that the user refer to NCCLS EP-12A⁸ and 42 CFR 493.1202(c) for guidance on appropriate QC practices. The external positive and negative controls are intended to monitor substantial reagent failure. The positive control will not challenge the assay at the cutoff.

If the controls do not perform as expected, repeat the test or contact Wampole Technical Support before testing patient specimens. **If the controls do not work as expected, a report should not be generated to the clinician.**



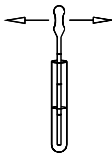
TEST PROCEDURES

Expiration date: Check expiration date on each individual test package (foil package or outer box) before using. *Do not use any test past the expiration date on the label.*



Nasopharyngeal swab procedure

1. Vortex or agitate the swab and transport medium or saline solution for 15-20 seconds to dislodge specimen from the swab.
2. Roll the swab head against the inside of the transport tube or specimen container as you remove it. Dispose of the used swab in accordance with your biohazard waste disposal protocol.



3. Draw 160 uL (second notch) of the sample up into a Disposable Pipette.



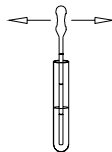
4. Add entire contents of the Pipette to the Extraction Tube.



5. Add four (4) **free falling** drops of R1 to the Extraction Tube while holding the bottle vertically over the Tube.



6. Gently swirl the Extraction Tube to mix the sample and R1.



7. Draw entire contents of the Extraction Tube up into a clean Disposable Pipette.



8. Squeeze entire contents of the Pipette drop wise onto the sample well of the Test Device.

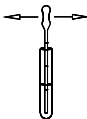


9. The test result should be read between fifteen (15) and twenty-five (25) minutes after applying the extracted sample to the Test Device. Results read outside the recommended time range are considered invalid.

Nasal Aspirate Procedure



1. Gently agitate with swirling motion or vortex aspirate sample to suspend cellular material in sample.



2. Fill the Disposable Pipette to the second notch (160uL) with nasal aspirate sample.



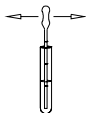
3. Add entire contents of the Disposable Pipette to the Extraction Tube.



4. Add four (4) **free falling** drops of R1 to the Extraction Tube while holding the bottle vertically over the tube



5. Gently swirl the Extraction Tube to mix the sample and R1.



6. Draw entire contents of Extraction Tube up into a clean Disposable Pipette.



7. Squeeze entire contents of the Pipette drop wise onto the sample well of the Test Device.



8. The test result should be read between fifteen (15) and twenty-five (25) minutes after applying the extracted sample to the Test Device. Results read outside the recommended time range are considered invalid.

Positive Result:

For a positive specimen, the appearance of **TWO** pink to red lines, one at the procedural Control Line (C) and one at the Test Line (T) indicates the presence of respiratory syncytial virus antigen. Any pink to red test line, even if it is only slightly pink, is considered a positive test. If the results are read after 25 minutes, the result is invalid. Specimen should be retested. **Results reported:** Positive for the presence of RSV antigen. A positive result may occur in the presence of both viable and non-viable virus.

Negative Result:

For a negative specimen, the appearance of **ONLY ONE** pink to red line at the procedural Control Line (C) and no pink to red line at the Test Line (T) indicates that the sample is negative for RSV viral antigen. If results are read after 25 minutes, the result is invalid. Specimen should be retested. **Results reported:** Negative for presence of RSV antigen. Infection due to RSV cannot be ruled out since the antigen present in the sample may be below the detection limit of the test. Cell Culture or DFA confirmation of negative samples is recommended.

Invalid Result:

If after 15 minutes, the pink to red procedural Control Line (C) does not appear, even if any shade of pink to red Test Line (T) appears, the result is considered invalid. If the test is considered invalid because a control line fails to appear, the test should be repeated with a new Test Device.

Within 15 minutes the result area should be white to light pink and allow the clear interpretation of the test result. **If the background color persists and interferes with the interpretation of the test result, the result is considered invalid.** Should this occur, review the test procedure and repeat the test with a new Test Device.

If the test result is still invalid after repeating with a new Test Device, then it is considered invalid.

If results are invalid, a report should not be generated to the clinician

LIMITATIONS

See Table 8 for limits of detection of Clearview RSV.

The contents of this kit are to be used for the qualitative detection of respiratory syncytial virus antigen from nasopharyngeal swab and nasal aspirate specimens.

Failure to follow the Test Procedure or Interpretation of Test Results instructions may adversely affect test performance and/or invalidate the test result.

Test results must be evaluated in conjunction with other clinical data available to the physician.

A negative test result may occur if the level of antigen in a sample is below the detection limit of the test, or from improper sample collection and storage.

Fresh specimens are preferable to frozen for RSV testing because of the highly thermolabile and fragile nature of the virus. Sub-optimal test performance may result with the latter.

A negative test result does not rule out other microbial respiratory tract infections.

Cross-reactivity of this assay with human metapneumovirus has not been studied.

Monoclonal antibodies may not detect all antigenic variants or new strains of RSV.

Testing with both the sample and/or the kit should not be performed at 4°C. Kit performance could be diminished if either the sample and/or the kit are not brought to room temperature first.

Testing of low levels of RSV in high humidity at room temperature could reduce the test's ability to detect RSV.

Testing with total sample volumes less than or greater than the recommended volume could reduce the ability of the test to detect low levels of RSV.

Low level positive results may not be properly visualized at ≤30% 'normal' bench lighting levels.

EXPECTED VALUES

The rate of positivity observed in RSV testing will vary depending on the time of year, age of the patient, geographic location, and local disease prevalence¹. In temperate climates, RSV infections occur primarily during annual outbreaks which peak during the winter months⁴. RSV is a common pathogen among infants and young children⁵. However, it may cause serious lower respiratory tract illness throughout life, especially among the elderly and those with compromised immune systems⁶⁻⁷. The following table summarizes the age and sex demographic data with the number of samples in each age group and the percent positivity for that group among the clinical samples tested by Clearview RSV. There are a total of 180 patients whose age was given with the sample included in the table. Of these, 92 had the sex of the patient recorded with the sample. See Table 1 below

Table 1: Patient Demographics

Patient Age	M:F	N	% Positive Clearview Results
2d-1 M	2:2	12	5/12 = 41.6%
1-2 M	2:2	8	4/8 = 50%
2M- 6 Y	22:12	112	30/112 =26.7%
51-60 Y	1:1	2	0/2 = 0%
61-70 Y	2:7	9	3/9 = 33.3%
71-80 Y	1:6	7	0/7 = 0%
81-90 Y	7:6	13	2/13 = 15.5%
91-100 Y	6:11	17	2/17 = 11.7%

N = total number of patient samples in each age cohort

Of the 17 patients whose age was not recorded, 68.7% were positive by Clearview RSV and 62.5% were positive by the reference method. Nine patients whose age was recorded as “greater than 60” were” were placed in the 61-70 year age group.

CLIA CONSIDERATIONS

This is a waived test under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) so long as it is used according to the instructions set in this package insert.

Any modification by the laboratory to the test system or the test system instructions will result in this test no longer meeting the requirements for waived categorization. A modified test is considered high complexity and is subject to applicable CLIA requirements. Further, the laboratory should notify Wampole Laboratories of any performance, perceived or validated, that does not meet the performance specifications as outlined in the instructions.

Under CLIA, Consumer Precision Studies and Consumer Accuracy Studies were conducted to demonstrate that lay users with no formal laboratory training could read the package insert instructions and perform the test with a high level of concordance with trained laboratorians. Consumer Precision Study testing was conducted using 90 swab samples at negative (0 pfu/mL RSV), LOD (1.9×10^5 pfu/mL RSV), and moderate positive (commercial kit positive control at 3.8×10^5 pfu/mL RSV) levels.

No significant differences were observed between the lay users and the trained laboratorians (Consumer Accuracy Study) or between the lay users within each site and among the three sites (Consumer Precision Study).

A summary table of the Consumer Accuracy Study and Consumer Precision Study is shown below (Table 2).

Table 2: CLIA Consumer Precision/Accuracy Studies

Participants	Frequencies of Test Results With 95% C.I.					
	Negative		LOD		Mod Positive	
	% + Test	% - Test	% + Test	% - Test	% + Test	% - Test
Lay Users (3 Sites)	0% 0/90	100% 90/90	94.4% 85/90	5.5% 5/90	100% 90/90	0% 0/90
	(0.0 - 4.1%)	(95.9 - 100%)	(87.6 - 97.6%)	(2.4 - 12.4%)	(95.9 - 100%)	(0.0 - 4.1%)
Laboratorian (3 sites)	0% 0/90	100% 90/90	96.7% 87/90	3.3% 3/90	100% 90/90	0% 0/90
	(0.0 - 4.1%)	(95.9 - 100%)	(90.7 - 98.9%)	(1.1 - 9.3%)	(95.9 - 100%)	(0.0 - 4.1%)

PERFORMANCE CHARACTERISTICS

The performance of the Clearview RSV Test was compared to Tissue Culture, EIA and Direct Fluorescent Antibody (DFA) in a prospective multi-center field clinical study in the Midwest and Ontario Canada during the 2002 flu season. In this trial, the Clearview RSV test was compared to DFA at the two United States sites or Tissue Culture and EIA at one Canadian site. The study comprised a total of 197 nasal aspirate and nasopharyngeal swab samples obtained from patients symptomatic for RSV infection. All samples were split to allow testing of the same sample by Clearview and the reference method (Tissue Culture and EIA at the Canadian site or DFA at the U.S. sites). The age of the patients ranged from 2 days to 100 years of age. The sample type in the U.S. consisted of nasopharyngeal swabs and aspirates and the sample type in Canada was entirely nasopharyngeal swabs.

United States Data

There were a **total of 104** samples tested in the United States. There were **90** nasal aspirates, **7** nasopharyngeal swabs, and **7** samples with no designation of sample type given. All 104 of the samples tested in the United States were compared to DFA as the reference method as shown in **Table 3**

Table 3: United States: Clearview RSV Compared to DFA

		DFA Result	
		Pos	Neg
Clearview Results	Pos	42	2
	Neg	3	57

Pos Agreement: 42/45 = 93.3% (95% C.I. = 80.7 -98.6%)

Neg Agreement: 57/59 = 96.6% (95% C.I. = 88.3 -99.6%)

A subset of the U.S data, the **90** samples designated as nasal aspirates is shown in **Table 4**.

Table 4: United States: Clearview RSV Nasal Aspirate Samples Compared to DFA

		DFA Result	
		Pos	Neg
Clearview Results	Pos	35	2
	Neg	2	51

Pos Agreement: 35/37= 94.6% (95% C.I. = 81.8- 99.3%)

Neg Agreement: 51/53= 96.2% (95% C.I. = 87.7- 99.3%)

Note: The 7 samples with no sample type designation and the 7 nasopharyngeal swab samples in the U.S. have not been broken out into separate tables but are included in Table 2. Of the 7 samples designated as nasopharyngeal swabs in the U.S., 3 were positive and 4 were negative and there was 100% positive agreement and 100% negative agreement with DFA.

Canadian Data

There were a **total of 93** samples tested in Canada. All were nasopharyngeal swab samples. The reference test for **92** of these was **both** Tissue Culture and EIA and **one** sample had **only EIA** as the reference for a total of **93**. The performance data for Clearview against the two reference methods is shown in **Tables 5 and 6**.

Table 5: Clearview RSV Compared to Tissue Culture in Canada (All Nasopharyngeal Swabs)

		Tissue Culture Result	
		Pos	Neg
Clearview Results	Pos	10	10
	Neg	0	72

Sensitivity: 10/10 = 100% (95% C.I. = 69.2 – 100 %)

Specificity: 72/82 = 87.8% (95% C.I. = 80.7 – 94.9%)

Table 6: Clearview RSV Compared to EIA in Canada (All Nasopharyngeal Swabs)

		EIA Result	
		Pos	Neg
Clearview Results	Pos	20	1
	Neg	1	71

Positive Agreement: 20/21= 95.2% (95% C.I. = 76.2- 99.9%)

Negative Agreement: 71/72=96.6% (95% C.I. = 92.5- 99.9%)

Summary of Clinical Data For All Sites

For all samples at all sites: There were a total of **197** samples, **104** in the U.S. and **93** in Canada:

Overall positive agreement = 93.9% (95% C.I. = 85.4-97.6%)

Overall negative agreement = 97.7% (95% C.I. = 93.5-99.2%)

For Nasopharyngeal Swabs: There were a total of **100** samples, **7** in the U.S. and **93** in Canada:

Overall positive agreement = 95.8% (95% C.I. = 79.8 – 99.3%)

Overall negative agreement= 98.7% (95% C.I. = 92.9 – 99.8%)

For Nasal Aspirates There were a total of **90** samples in the U.S.

Positive Agreement: = 94.6% (95% C.I. = 81.8 - **99.3%**)

Negative Agreement: = 96.2% (95% C.I. = 87.7 - **99.3%**)

Note: 7 samples in the U.S. had no designation of sample type and are not included in the breakdown of nasopharyngeal swab performance (N = 100) and nasal aspirate performance (N = 90). These 7 samples are included in the overall performance data for all samples at all sites (N = 197) and in the table for all of the U.S. data (Table 2).

ANALYTICAL SPECIFICITY AND CROSS-REACTIVITY

The bacterial and viral organisms listed below were used to assess cross reactivity in the Clearview RSV Test. No cross reactivity was noted with any of the organisms tested. The testing was done in replicates of three. (See **Table 7 and 8**)

Table 7: Bacterial Panel

Organism Tested	Strain/Reference	Final Concentration On Test Device	Results	
			RSV Interference	Microbe
<i>Candida Albicans</i>	10231	5.0 x 10 ⁶ org/mL**	3/3 pos	3/3 neg
<i>Chlamydia psittaci</i>	VMLCPS0427	2.0 x 10 ⁷ inc/mL***	3/3 pos	3/3 neg
<i>Chlamydia trachomatis</i>	VML CT 092801	2.0 x 10 ⁶ inc/mL	3/3 pos	3/3 neg
<i>Haemophilus influenzae</i>	10211	5.0 x 10 ⁷ org/ml	3/3 pos	3/3 neg
<i>Klebsiella pneumoniae</i>	13883	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Mycoplasma pneumoniae</i>	15531	2.6 x 10 ⁷ CFU/ml *	3/3 pos	3/3 neg
<i>Neisseria meningitidis</i>	13090	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Pseudomonas aeruginosa</i>	9027	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Serratia liquifans</i>	27592	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Staphylococcus aureus</i>	25923	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Staphylococcus aureus</i>	Cowan	3.0 x 10 ⁸ org/mL	3/3 pos	3/3 neg
<i>Staph epidermidis</i>	12228	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Strep pneumoniae</i>	Wild strain	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Streptococcus sp gr A</i>	12384	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Streptococcus sp gr F</i>	12392	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg
<i>Streptococcus sp gr G</i>	12394	5.0 x 10 ⁷ org/mL	3/3 pos	3/3 neg

***The Cowan Strain of Staphylococcus aureus is a Protein A producing strain and did not show cross reactivity with Clearview RSV.**

****org/mL = organisms/mL**

*****inc/mL = inclusions/mL**

Table 8: Viral Panel

Test Organism	Strain/ Reference	Final Concentration On Test Device	Results	
			Clearview RSV Interference	Microbe
<i>Adenovirus 5</i>	VR-5	1.0 x 10 ⁵ PFU/mL	3/3 pos	3/3 neg
<i>Adenovirus 7</i>	VR-7	1.0 x 10 ⁵ PFU/mL	3/3 pos	3/3 neg
<i>Adenovirus 2</i>	VR-846	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Coxsackie A19</i>	VR-1015	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Coxsackie B1</i>	VR- 28	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Coxsackie B3</i>	VR-30	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Cytomegalovirus</i>	AD-169	8.8 x 10 ⁴ PFU/mL	3/3 pos	3/3 neg
<i>Poliovirus</i>	VR-1004	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Echovirus 6</i>	VR-36	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Echovirus 11</i>	VR-41	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>HSV Type 1</i>	VR-6143	3.3 x 10 ⁵ PFU/mL	3/3 pos	3/3 neg
<i>HSV Type 2</i>	VR-15671	3.3 x 10 ⁵ PFU/mL	3/3 pos	3/3 neg
<i>Influenza A</i>	N1H1	9.2 x 10 ² PFU/mL	3/3 pos	3/3 neg
<i>Influenza B</i>	Beijing	5.9 x 10 ³ PFU/mL	3/3 pos	3/3 neg
<i>Parainfluenza 1</i>	VR-105	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Parainfluenza 2</i>	VR-92	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Parainfluenza 3</i>	VR-93	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Rhinovirus 1B</i>	VR-1366	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Rhinovirus 39</i>	VR-340	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Rhinovirus 16</i>	VR-1126	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Rhinovirus 14</i>	VR-284	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Rhinovirus 37</i>	VR-1147	1.0 x 10 ⁵ TCID ₅₀ /mL	3/3 pos	3/3 neg
<i>Varicella</i>	Ellen	6.5 x 10 ³ PFU/mL*	3/3 pos	3/3 neg

These viral strains were obtained from American Type Culture Collection (ATCC) and the titers were established by the independent laboratory where cross reactivity testing was performed.

ANALYTICAL SENSITIVITY

The Clearview RSV strip test limit of detection (LOD) for RSV antigen subgroup A (Long and A2 strains) and antigen subgroup B (18537 and WV/14617/85, B-1 strains) are shown in **Table 9**.

Table 9: Clearview RSV Limits of Detection (LOD)

RSV Strain	Antigen Subgroup	LOD pfu/mL	Frequency Positive Test	95% Confidence Interval
Long	A	1.9X10 ⁵	90%	68.3-98.7%
A2	A	>1.12 x 10 ⁵	80%	56.0-94.2%
18537	B	>1.9X10 ⁵	80%	56.0-94.2%
WV14617/85	B	1.9X10 ⁵	95%	75.0-99.9%

All concentrations of RSV are those found in original samples before manipulation and placement on the Clearview RSV test device.

INTERFERING SUBSTANCES

Whole blood, mucin, and several over the counter (OTC) products, common chemicals, storage media, and transport media were evaluated and did not interfere with the Clearview RSV Test at the levels tested. The testing was done in replicates of three. (See **Table 10**)

Table 10: Interfering Substances Panel

Test Substance	Final Conc. On Test Device	Results	
		RSV Competition	Interfering Substance
Scope Mouthwash	25%	3/3 positive	3/3 negative
Listerine Mouthwash	25%	3/3 positive	3/3 negative
CVS Mouthwash	25%	3/3 positive	3/3 negative
Ludens Cough Drops	25%	3/3 positive	3/3 negative
Hall Cough Drops	25%	3/3 positive	3/3 negative
Ricola Cough Drops	25%	3/3 positive	3/3 negative
Afrin Nasal Spray	10%	3/3 positive	3/3 negative
Neosynephrine N S	10%	3/3 positive	3/3 negative
CVS Nasal Spray	10%	3/3 positive	3/3 negative
4-Acetamidophenol	10 mg/mL	3/3 positive	3/3 negative
Acetylsalicylic Acid	20 mg/mL	3/3 positive	3/3 negative
Chlorpheniramine	5 mg/mL	3/3 positive	3/3 negative
Dextromethorphan	10 mg/mL	3/3 positive	3/3 negative
Diphenhydramine	5 mg/mL	3/3 positive	3/3 negative
Ephedrine	20 mg/mL	3/3 positive	3/3 negative
Guaiacol Glycerol Ether	20 mg/mL	3/3 positive	3/3 negative
Oxymetazoline	10 mg/mL	3/3 positive	3/3 negative
Albuterol	10 mg/mL	3/3 positive	3/3 negative
Mucin	4 mg/mL	3/3 positive	3/3 negative
Whole Blood	50% w/v	3/3 positive	3/3 negative

REPRODUCIBILITY STUDIES

NCCLS document EP 12, User Protocol for Evaluation of Qualitative Test Performance; Approved Guideline⁸ defines critical values for evaluation of performance of a qualitative test. These critical values include the “cutoff” or the test response above which the test result is determined to be positive and below which the test response is determined to be negative, and the LOD at which the test response gives 95% positive results and 5% negative results. The cutoff is the level of virus that gives 50% positive results and 50% negative results. Levels at 10% and 20% above the cutoff should give rates of positivity between 50% and 95%. For levels of RSV at 10% and 20% below the cutoff the rate of negative test results should be between 50% and 95%.

Over 3 consecutive days, at numerous test sites, and by multiple operators, the Clearview RSV test performance was evaluated by both in-house masked reproducibility trials (see **Table 11**) and physician office laboratory masked reproducibility trials (see **Table 12**). As shown, these trials define the values critical to qualitative function and include:

1. The cutoff at 9.3×10^4 pfu/mL of virus where 50% of replicate samples are found to be positive and 50% are found to be negative.
2. The limit of detection (LOD) at 1.9×10^5 pfu/mL of virus where positive test results approach 95% and negative test results approach 5%.
3. The region of virus concentration (pfu/mL) above the cutoff where positive test values range between 50% and 95%.
4. The region of virus concentration (pfu/mL) below the cutoff where negative test results range between 50% and 95% respectively.

Additionally, POL field studies at 3 independent sites showed expected results with values of percent positive test for the LOD at 1.9×10^5 pfu/mL that approach 95%. Levels of virus above the 95% interval at 7.7×10^5 pfu/mL that approach 100% positive test results are shown. Also, levels of virus at 10% (1.02×10^5 pfu/mL) and 20% (1.12×10^5 pfu/mL) above the cutoff have expected positive test results between 61% and 74% respectively. At 20% below the cutoff (7.4×10^4 pfu/mL) the rate of negative test results is 77% and outside the 95% interval at 4.7×10^4 pfu/mL the rate of negative test results approach 100% (**Table 12**).

Note: The positive control used in the reproducibility trials is the RSV Long Strain at the LOD (1.9×10^5 pfu/mL) of the device in order to better challenge this critical point of the assay. The control included in the kit is twice this LOD concentration.

In House Reproducibility

Table 11 below shows Clearview RSV positive and negative test results with 95% confidence intervals (C.I.) for 5 concentrations of RSV: (1) limit of detection (LOD) of 1.9×10^5 pfu/mL, (2) cutoff of 9.3×10^4 pfu/mL, (3) 20% above the cutoff at 1.12×10^5 pfu/mL, (4) 10% above the cutoff at 1.02×10^5 pfu/mL and, (5) 20% below the cutoff at 7.4×10^4 pfu/mL. Testing was done with coded samples as daily runs over 3 consecutive days by 3 operators. All concentrations of RSV are those found in original samples before manipulation and placement on the Clearview RSV test device.

Table 11: Clearview RSV In-House Reproducibility

RSV Long Strain (pfu/mL)	Number Replicates	Frequency (%) Positives (95% C.I.)	Frequency (%) Negatives (95% C.I.)
1.9×10^5	30	83.3 (65.3 – 94.4)	16.7 (5.64 – 34.7)
1.12×10^5	90	67.8 (57.1 – 77.3)	32.2 (28.8 – 42.9)
1.02×10^5	90	58.9 (48.0 – 69.2)	41.1 (30.8 – 51.9)
9.30×10^4	90	50.0 (39.3 – 60.7)	50.0 (39.3 – 60.7)
7.40×10^4	90	23.3 (15.1 – 33.4)	76.6 (66.6 – 84.4)

Physician Office Laboratory (POL) Reproducibility Studies

Table 12 shows Clearview RSV positive and negative test results with 95% confidence intervals (C.I.) for concentrations of RSV at the limit of detection (LOD) of 1.9×10^5 pfu/mL, above the 95% interval at 7.7×10^5 pfu/mL, three levels around the cutoff ranging from 7.40×10^4 pfu/mL to 1.12×10^5 pfu/mL and, below the 95% interval at 4.7×10^4 pfu/mL. Testing was done with coded samples by 6 operators with diverse educational background and work experiences in replicates of 3 at 3 different locations over 3 days. All concentrations of RSV are those found in original samples before manipulation and placement on the Clearview RSV test device.

Table 12: Clearview RSV POL Reproducibility

RSV Long Strain (pfu/mL)	Number Replicates	Frequency (%) Positives (95% C.I.)	Frequency (%) Negatives (95% C.I.)
7.7 x 10 ⁵	54	100 (93.4 - 100)	00.0 (0.0 – 6.6)
1.9 x 10 ⁵	54	100 (93.4 - 100)	00.0 (0.0 – 6.6)
1.12 x 10 ⁵	54	74.1 (60.4 - 85.0)	25.9 (14.9 – 39.65)
1.02 x 10 ⁵	54	61.1 (46.9 – 74.1)	38.8 (25.9 – 53.12)
7.40 x 10 ⁴	54	22.2 (12.0 – 77.7)	77.7 (64.4 – 87.96)
4.7 x 10 ⁴	54	00.0 (0.0 – 6.6)	100 (93.4 - 100)

Note: The RSV Long strain used to conduct these reproducibility studies is a laboratory adapted strain and may or may not perform the same as clinical specimens.

ASSISTANCE

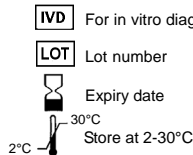
If you have any questions regarding the use of this product, please call Wampole's Technical Service at 1-800-257-9525 (press 2). If outside the United States, please contact your local distributor.

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