



ARTG ID: 524286

English

COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit Instruction for Use

INTENDED USE

The COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit is a lateral flow immunoassay for the qualitative detection of SARS-CoV-2 (nucleocapsid protein), Respiratory Syncytial Virus (nucleocapsid protein), Adenovirus (Hexon Protein), Influenza A (nucleocapsid protein) and Influenza B (nucleocapsid protein) in nasal swabs from subjects. The symptoms of respiratory viral infection due to SARS-CoV-2, respiratory syncytial, Adenovirus, influenza can be similar. The test is intended as an aid in diagnosis of symptomatic individual meeting respiratory infection for SARS-CoV-2 (within the first 7 days of the onset of symptoms) and influenza A/B, Adenovirus or Respiratory syncytial virus (RSV) (within the first 4 days of the onset of symptoms). This kit is intended for layperson's home use in a non-laboratory environment.

PRINCIPLE

The COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit is a lateral flow immunoassay based on the principle of the double antibody sandwich technique. A monoclonal SARS-CoV-2/RSV/ADV/Influenza A&B antibody conjugated with colored microparticles and sprayed onto the conjugation pad is used as a detector. During the test, the SARS-CoV-2/RSV/ADV/Influenza A&B antigen in the sample interacts with the SARS-CoV-2/RSV/ADV/Influenza A&B antibody conjugated with colored microparticles, creating an antigen-antibody labeled complex. This complex migrates on the membrane by capillary action up to the Test line where it is captured by the pre-coated monoclonal SARS-CoV-2/RSV/ADV/Influenza A&B antibodies. A colored test line (T) would be visible in the each result window if SARS-CoV-2/RSV/ADV/Influenza A&B antigens are present in the sample. The absence of the T line indicates a negative result. The control line (C) is for procedural control and should appear whenever the test procedure is being performed properly.

PRECAUTIONS

- 1. For in vitro diagnostic use only.
2. Do not use after the expiration date.
3. Perform the test at room temperature 15 to 30°C.
4. The test cassette should remain in the sealed pouch until use.
5. Please read all information in this leaflet before performing the test.
6. Components from difference lots must not be mixed or used together.
7. Positive result cannot necessarily determine whether a person is infectious.

STORAGE AND STABILITY

Store the test kit in the original packaging at 2°C - 30°C. Do not freeze. Test kit contents remain stable until the expiration date printed on the outer packaging. After opening the pouch, the test should be used within one hour. Prolonged contact with hot and humid environment will cause the product to deteriorate.

LIMITATION

- 1. A negative result does not rule out infection with another type of respiratory virus. And a positive result cannot necessarily determine whether a person is infectious.
2. The test can only be used once.
3. Test can only be performed by person over 15 years age. Any persons or children under 15 years will require adult supervision or assistance. Not to be performed on children under 2 years of age.
4. The performance of COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit was established based on the evaluation of a limited number of clinical specimens. Clinical performance has not been established with all circulating variants but is anticipated to be reflective of the prevalent variants in circulation at the time and location of the clinical evaluation. Performance at the time of testing may vary depending on the variants circulating, including newly emerging strains of SARS-CoV-2 and their prevalence, which change over time.
5. In particular, false negative results may occur if the testing is not performed within the first 7 days of the onset of COVID-19 or within the first 4 days of influenza A&B/Adenovirus/RSV symptoms or if the antigen level in the sample is below the detection limit.
6. The tests are less reliable in the later phase of infection and in asymptomatic individuals.
7. Recommend repeat testing within 1-3 days if there is an ongoing suspicion of infection, being in a high risk setting or where there is an occupational risk or other requirement.
8. Negative results may not mean that a person is not infectious and if symptoms persist, please seek medical advice.

SAFETY INFORMATION

- 1. Wearing a mask can help protect you and those around you if you are in an area with community transmission and physical distancing is not possible.
2. Follow the directions of your local state or territory government health department to protect yourself.
3. Test kit solutions should only be used as directed; do not ingest; do not dip the swab into provided solution or other liquid before inserting the swab into the nose; avoid contact with skin and eyes; keep out of the reach of children and pets before and after use.
4. If the extraction buffer comes in contact with the skin or eyes, flush with plenty of water. If irritation persists, seek medical advice from a doctor or your local medical center.

CLINICAL PERFORMANCE

For COVID-19
The clinical performance of the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit for professional use has been evaluated against RT-PCR, and the results are as follows:
Clinical sensitivity = 97.08% (498/513 known confirmed Positives, 95%CI* 95.23% to 98.22%)
Clinical specificity = 99.07% (743/750 known confirmed Negatives, 95%CI* 98.09% to 99.55%)
*.95% confidence interval

Table with 4 columns: Days after symptom onset, RT-PCR, ACRT, Sensitivity. Rows show data for days 0-7 and stratified by age groups (0-5, 6-21, 22-59, ≥60).

For influenza A test

The clinical performance of the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit for professional use has been evaluated against RT-PCR, and the results are as follows:
Clinical sensitivity = 94.62% (123/130 known confirmed Positives, 95%CI* 89.30% to 97.37%)
Clinical specificity = 98.74% (157/159 known confirmed Negatives, 95%CI* 95.53% to 99.65%)
*.95% confidence interval

Two tables: Sensitivity Result according to the date of onset and Sensitivity Result stratified by age group. Rows show data for days after symptom onset and age groups.

For influenza B test

The clinical performance of the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit for professional use has been evaluated against RT-PCR, and the results are as follows:
Clinical sensitivity = 92.31% (120/130 known confirmed Positives, 95%CI* 86.42% to 95.77%)
Clinical specificity = 98.21% (165/168 known confirmed Negatives, 95%CI* 94.88% to 99.39%)
*.95% confidence interval

Two tables: Sensitivity Result according to the date of onset and Sensitivity Result stratified by age group. Rows show data for days after symptom onset and age groups.

For RSV test

The clinical performance of the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit for professional use has been evaluated against RT-PCR, and the results are as follows:
Clinical sensitivity = 95.56% (129/135 known confirmed Positives, 95%CI* 90.64% to 97.95%)
Clinical specificity = 99.16% (118/119 known confirmed Negatives, 95%CI* 95.39% to 99.85%)
*.95% confidence interval

Two tables: Sensitivity Result according to the date of onset and Sensitivity Result stratified by age group. Rows show data for days after symptom onset and age groups.

For Adenovirus test

The clinical performance of the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit for professional use has been evaluated against RT-PCR, and the results are as follows:
Clinical sensitivity = 95.83% (115/120 known confirmed Positives, 95%CI* 90.62% to 98.21%)
Clinical specificity = 98.36% (120/122 known confirmed Negatives, 95%CI* 94.22% to 99.55%)
*.95% confidence interval

Two tables: Sensitivity Result according to the date of onset and Sensitivity Result stratified by age group. Rows show data for days after symptom onset and age groups.

USABILITY STUDY RESULTS

The usability study was conducted with lay persons who performed the test and interpreted the result. The test procedure and obtain consistent test results with professionals.
For SARS-CoV-2, The results were compared to an RT-PCR with a sensitivity of 94.29% (33/35) and specificity of 100% (65/65).
For Influenza A, The results were compared to an RT-PCR with a sensitivity of 94.44% (34/36) and specificity of 98.44% (63/64).
For Influenza B, The results were compared to an RT-PCR with a sensitivity of 90.63% (29/32) and specificity of 98.53% (67/68).
For ADV, The results were compared to an RT-PCR with a sensitivity of 94.12% (32/34) and specificity of 98.48% (65/66).
For RSV, The results were compared to an RT-PCR with a sensitivity of 93.75% (30/32) and specificity of 98.53% (67/68).

LIMIT OF DETECTION (ANALYTICAL SENSITIVITY)

Table with 2 columns: Virus Lines, LoD Titer. Rows list various virus strains and their corresponding LoD titers.

Table with 2 columns: Virus Lineage, LoD Titer. Rows list various virus lineages and their corresponding LoD titers.

FREQUENTLY ASKED QUESTIONS

1. Will other diseases affect the result?

The potential cross-reactivity of the following pathogens was evaluated with SARS-CoV-2, respiratory syncytial, Adenovirus, Influenza A and B negative and positive samples using the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit. No interference or competitive inhibition was observed.

Table with 3 columns: Virus or organisms, SARS-CoV, Influenza A H5N1 virus, etc. Lists various viruses and their detection status.

2. Does these substances interfere with the test?

The following substances were spiked into samples and tested with the COVID-19/RSV/ADV/Flu A&B Antigen Combo Rapid Test Kit. No interference results was observed.

Table with 3 columns: Substances, Mucin, Human blood, etc. Lists various substances and their detection status.

3. Will this test hurt?

No, the nasal swab is not sharp and it should not hurt. Sometimes the swab can feel slightly uncomfortable or tickly. If you feel pain, please stop the test and seek advice from a healthcare provider.

4. I have a nosebleed after swabbing my nose. What should I do?

In the unlikely event your nose starts bleeding, apply pressure to your nose until the bleeding stops and consult a healthcare professional. Do not insert the Swab again.

5. Can the test detect various variants of COVID-19?

Yes, the following SARS-CoV-2 variants can be detected with the COVID-19 (SARS-CoV-2) Antigen Test Kit: Alpha, Beta, Gamma, Delta and Omicron.

6. Which strains of influenza the test covers?

Table with 3 columns: Influenza A, A/Vietnam/HN31242/2007, A/Shanghai/2/2013, etc. Lists various influenza A strains.

Table with 3 columns: Influenza B, B/Sichuan/Gaoxin/531/2018, B/Hong Kong/3417/2014, etc. Lists various influenza B strains.

7. Which strains of RSV the test covers?

Table with 3 columns: RSV Type A/A2, RSV Type A/Long, RSV Type B/GZ/1704-8, etc. Lists various RSV strains.

8. Which strains of Adenovirus the test cover?

Table with 3 columns: Adenovirus 1/GZ/1608-21, Adenovirus 2/GZ/1705-34, Adenovirus 3/GZ/0101/2011, etc. Lists various Adenovirus strains.

MEDICAL DEVICE INCIDENT REPORT

You can contact the Therapeutic Goods Administration (TGA) to report poor performance or usability issues via the online Users Medical Device Incident Report, emailing iris@health.gov.au or calling 1800 809 361 (08:30am to 5:00pm Monday to Friday).

Note: Use test only one time.

Scan the QR code or visit our website for instructional video, product information and IFU:
<https://www.ahpcpharmacyoutlet.com.au/products/cellife-5-in-1-antigen-test-kit-single-pack-pending-tga>

| Components | REF | ICRAF -555-E010 | ICRAF -555-E040 | ICRAF -555-E050 | ICRAF -555-E070 | ICRAF -555-E100 | ICRAF -555-E250 |
|---------------------------|-----|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1. Test Cassette | 1 x | 4 x | 5 x | 7 x | 10 x | 25 x | |
| 2. Extraction Buffer Tube | 1 x | 4 x | 5 x | 7 x | 10 x | 25 x | |
| 3. Disposable Swab | 1 x | 4 x | 5 x | 7 x | 10 x | 25 x | |
| 4. Instruction for Use | 1 x | 1 x | 1 x | 1 x | 1 x | 25 x | |

Materials required but not provided : Timer

For the sterilized swab
 CE 0197 MDR 2017/745 EU Hangzhou Yiguoeren Biotechnology Co., Ltd.
 CE 0197 MDD 95/42/EEC Jiangsu Han-Heng Medical Technology Co., Ltd.

COVID-19 / RSV / ADV

Influenza A&B

NOTE

The intensity of color that the test line area (COVID-19/Flu A/Flu B/ADV/RSV) shows will vary according to the concentration of SARS-CoV-2 antigen, Influenza A antigen and Influenza B antigen, ADV antigen, RSV antigen. The result should be determined on whether the test line is formed or not, and is irrelevant to the color intensity. Therefore, any intensity of color in the test area (COVID-19/Flu A/Flu B/ADV/RSV) should be considered positive.

Positive result:

- SARS-CoV-2 positive result:** If both the control line (C line) and the test line (T line) appear at the same time, it means that SARS-CoV-2 antigen has been detected and the result is positive. There is currently a suspicion of a COVID-19 infection.
- If you test positive, you should not visit high-risk settings like hospitals and aged and disability care settings for at least 7 days or until symptoms have gone, unless seeking immediate medical care.
- To help protect those around you, we recommend to avoiding contact with people who are at higher risk of severe disease, wearing a mask outside the home, working from home where possible, avoiding going to school, public areas, or travel on public transport, in taxis or ride-share services, practicing good hygiene, and following your local health department's advice when leaving home.
- If you have any appointments you cannot miss (visit to a doctor, family violence service or police), let them know in advance that you have COVID-19.
- If you feel unwell or need COVID-19 advice for someone in your care, talk with your health provider, or speak to a nurse by calling the health direct helpline on 1800 022 222.
- If you develop symptoms such as severe shortness of breath or chest pain, call triple zero (000) immediately. Tell the call handler and the paramedics on arrival if you have COVID-19.

Influenza A positive result: If both the control line (C line) and the Influenza A test line (Flu A line) appear at the same time, it means that Influenza A antigen has been detected in the sample and the result of Influenza A is positive.

Influenza B positive result : If both the control line (C line) and the Influenza B test line (Flu B line) appear at the same time, it means that Influenza B antigen has been detected in the sample and the result of Influenza B is positive.

ADV positive result: If both the control line (C line) and the test line (T line) appear at the same time, it means that ADV antigen has been detected in the sample and the result of ADV is positive.

RSV positive result: If both the control line (C line) and the test line (T line) appear at the same time, it means that RSV antigen has been detected in the sample and the result of RSV is positive.

If the quality control C line appears, and more red lines appear in the detection line area, indicating that the sample contains one or more pathogenic microorganisms.

Negative result: If only the quality control C line appears and the detection line is not visible, the sample contains no SARS-CoV-2, Influenza A, Influenza B, ADV, RSV or the concentration is lower than the limit of detection and the result is negative.

Invalid result: If the C line does not appear, the result is invalid and a new test must be performed again.

Customer Support help line:
02 8054 5535
Customer Service hours:
9 AM ~ 8 PM, 7 Days. ~ UTC+10

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| | Do not re-use | | Use-by date |
| | In vitro diagnostic medical device | | Keep away from sunlight |
| | Store between 2-30°C | | Keep dry |
| | Consult instructions for use | | Do not use if package is damaged and consult instructions for use |
| | Batch code | | Manufacturer |
| | Contains sufficient for <n> tests | | Catalogue number |

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