

ARTG ID: 414738 REF: ICF - 535

English

COVID-19 / Influenza A&B Antigen Test Kit

INTENDED USE

The COVID-19/Influenza A&B Antigen Test Kit is a lateral flow immunoassay for the qualitative detection of SARS-CoV-2, influenza A and influenza B viral nucleoprotein antigens in nasal swabs. The symptoms of respiratory viral infection due to SARS-CoV-2 and influenza can be similar. The test is intended as an aid in diagnosis of symptomatic individual meeting the case definition for COVID-19 within the first 7 days of symptom onset and meeting the case definition for Influenza A&B within the first 4 days of symptoms onset.

This kit is intended for self-testing by laypersons in the home or other non-laboratory environment.

PRINCIPLE

The COVID-19/Influenza A&B Antigen Test Kit is qualitative detection of SARS-CoV-2, Influenza A and B antigens based on principal of immunochromatography. During testing, antigen in the specimen reacts with anti-SARS-CoV-2 antibody-coated particles and with anti-Influenza A antibody-coated particles as well as with anti-Influenza B antibody-coated particles in the conjugation pad to produce the immune complex. The complex migrates along the membrane by capillary action to the test region. The complex then respectively reacts with antibodies. If the specimen does contain antigen of SARS-CoV-2, Influenza A/B, colored line will appear in the test region, inflicating a positive result.

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.
- To avoid inaccurate results, use only the components in this test kit. Do not substitute test kit components from a different batch
- 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. 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 symptoms or if the antigen level in the sample is below the detection limit.
- $2. \ The \ tests \ are \ less \ reliable \ in \ the \ later \ phase \ of \ infection \ and \ in \ asymptomatic \ individuals.$
- Repeat antigen rapid testing is recommended every 24 hours for 3 days if there is a suspicion of infection, exposure to high-risk settings or other occupational risk.
- 4. 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.
- 5. The test can only be used once.
- 6. 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.

SAFETY INFORMATION

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.

Follow the directions of your local state or territory government health department to protect yourself.

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. 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 centre.

CLINICAL PERFORMANCE

For COVID-19

Using COVID-19/Influenza A&B Antigen Test Kit by professional was compared to the RT-PCR kit. A sensitivity of 92.5% (148/160 known confirmed Positives) and a specificity of 98.33% (295/300 known confirmed Negatives) were determined for the COVID-19 (SARS-CoV-2) Antigen Test Kit.

For influenza A tes

Using COVID-19/Influenza A&B Antigen Test Kit by professional was compared to the RT-PCR kit. A sensitivity of 88.12% (131/147 known confirmed Positives)and a Specificity of 98.33% (472/480 known confirmed Negatives) were determined for the COVID-19/Influenza A&B Antigen Test Kit.

For influenza B test

Using COVID-19/Influenza A&B Antigen Test Kit by professional was compared to the RT-PCR kit. A sensitivity of 89.86%(124/138 known confirmed Positives)and a Specificity of 98.18%(540/550 known confirmed Negatives)were determined for the COVID-19/Influenza A&B Antigen Test Kit.

Usability Study

110 lay users in different age distribution, different education level and different gender participated in usability study conducted in the self-testing environment. Compared to RT-PCR, the clinical performance of COVID-19/Influenza A+B Antigen Test kit in hands of lay persons showed a sensitivity of 92.3% (95% confidence interval: 79.68%-97.35%, N=39) and a specificity of 97.18% (95% confidence interval: 90.30%-99.22%, N=71) for COVID-19 antigen, a sensitivity of 87.5% (95% confidence interval: 73.89%-94.54%, N=40) and a specificity of 97.14% (95% confidence interval: 79.92.1%, N=70) for influenza A antigen and a sensitivity of 90.0% (95% confidence interval: 76.95%-96.04%, N=40) and a specificity of 95.71% (95% confidence interval: 88.14%-98.53%, N=70) for influenza B antigen.

LIMIT OF DETECTION (ANALYTICAL SENSITIVITY) LoD Titer (TCID so/ml.) SARS-CoV-2 wild type 1.0×10² 2.08×10³ Flu A H1N1/Wisconsin/588/2019 Flu A H3N2/SouthAustralia/34/2019 7.76×10² Flu B Austria/1359417/2021(Victoria lineage 2.84×103 Flu B Phuket/3073/2013 (Yamagata lineage) 1.08×104 3.105×10² Flu A H1N1/Reijng/262/95 Flu A H3N2/Shangdong/9/93 2.26×10² Flu B Victoria lineage/Shandong/7/97 1 825×103 Flu B Yamagata lineage/Jiangsu/10/03 2 44×103

FREQUENTLY ASKED QUESTIONS

1.Will other diseases affect the result?

The potential cross-reactivity of the following pathogens was evaluated with SARS-CoV-2, Influenza A and B negative and positive samples using the COVID-19/Influenza A&B Antigen Test Kit. No cross-reactivity results was observed.

Virus or organisms			
Human coronavirus NL63	Influenza A H5N1 virus	Coxsackie virus CA16e	
Human coronavirus HKU1	Influenza B Yamagata	Coxsackie virus B5	
Human coronavirus Oc43	Influenza B Victoria	Coxsackie virus A24	
Human coronavirus 229E	Haemophilus influenzae	Candida albicans	
MERS-CoV	Adenovirus 1	Human Metapneumovirus A2	
Respiratory syncytial virus Type A	Adenovirus 2	Legionella pneumophila	
Respiratory syncytial virus Type B	Adenovirus 3	Mycobacterium tuberculosis	
Parainfluenza virus 1	Adenovirus 4	Mycoplasma pneumoniae	
Parainfluenza virus 2	Adenovirus 5	Pneumocystis jiroveci	
Parainfluenza virus 3	Adenovirus 7	Streptococcus pneumoniae	
Parainfluenza virus 4	Adenovirus 55	Staphylococcus aureus	
Seasonal influenza A H1N1 virus	Enterovirus EV70	Rhinovirus A2	
Influenza A H3N2 virus	Bordete∎a pertussis	Rhinovirus B52	
SARS-CoV-1	Chlamydia pneumoniae	Streptococcus pyogenes	

2.Does these substances interfere with the test?

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

Substances				
Mucin	Phenylephrine Hydrochloride	Histamine hydrochloride		
Human blood (EDTA anticoagulated)	Arbidol	Alpha interferon		
Bedomethasone dipropionate nasal aerosol	Zanamivir	Azithromycin		
physiological seawater nasal spray	Ribavirin	Oseltamivir phosphate		
Triamcinolone acetonide nasal spray	Peramivir	Meropenem		
Mometasone furoate nasal spray	Lopinavir	Tobramycin		
Fluticasone propionate nasal spray	Ritonavir	Hexadecadrol		
Budesonide nasal spray	Levofloxacin	Flunisolide		
Oxymetazoline hydrochloride spray	Ceftriaxone			

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 ?

Influenza A				
A/Vietnam/HN31242/2007	A/Victoria/2570/2019	A/Brisbane/02/2018		
A/Shanghai/2/2013	A/Switzerland/8060/2017 A/Michigan/45/2015			
A/RR/8/34	A/Hong Kong/45/2019	A/Hong Kong/2671/2019		
A/California/04/2009	A/Wisconsin/588/2019			
A/Darwin/9/2021	A/Darwin/6/2021			
A/SouthAustralia/34/2019	A/Bean Goose/Hubei/chenhu XVI35-1/2016			
A/Guizhou/54/89	A/Singapore/INFMH-16-0019/2016			
	Influenza B			
B/SichuanGaoxin/531/2018	B/Austria/1359417/2021	B/Brisbane/60/2008		
B/Hong Kong/3417/2014	B/Washington/02/2019			
B/Phuket/3073/2013	B/Colorado/06/2017			

7. What to do if you test positive?

If you have a Covid-19 POSITIVE result, staying at home protects the people in your community and you should not visit high-risk settings like hospitals and aged and disability care settings.

If you feel unwell or need COVID-19 advice for someone in yourcare, talk with your health provider, or speak to a nurse by calling the healthdirect 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.

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).

SYMBOLS					
2	Do not re-use	\subseteq	Use-by date		
IVD	In vitro diagnostic medical device	巻	Keep away from sunlight		
2°C - 30°C	Store between 2-30°C	Ť	Keep dry		
	Consult instructions for use	8	Do not use if package is damaged and consult instructions for use		
LOT	Batch code	1	Manufacturer		
\sum	Contains sufficient for <n> tests</n>	REF	Catalogue number		

W

Hangzhou Fanttest Biotech Co.,Ltd.

Room 201, Building 1, No. 37-3, Futang Road, Tangqi Town, Linping District, Hangzhou City, Zhejiang Province,311106,P.R.China. E-mail: info@fanttest.com Tel: +86 571 86337555

Australia Sponsor

Australia Health Products Central Pty Ltd 604 / 3 Waverley St Bondi Junction Sydney NSW 2022

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

Web: www.cellifehealthcare.com.au

Code: 1054023000 Version No.: 1.0 Effective Date: Nov.11,2024



COVID-19 / Influenza A&B Antigen Test Kit

Note: Use test only one time.









tip with finge







Remove the swab from the LEFT nostril and insert into the RIGHT nostril. Again, rotate 5 times.



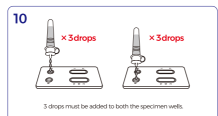
Remove the swab while squeezing the sides of the tube to extract the liquid from the swah



Scan the QR code or visit our website for instructional video, product information and IFU: https://www.ahpcpharmacyoutlet.com.au/ products/cellife-covid-19-influenza-a-bantigen-test-kit-single-pack











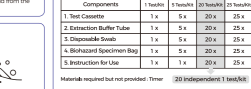
the walls of the tube 5 times. Allow

the swab to stand in the extraction

buffer tube for 1 minute

6

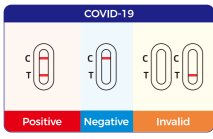


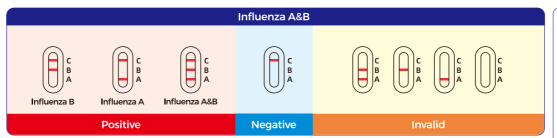




For the sterilized swab

CE 0197 MDR 2017/745 EU Hangzhou Yiguoren Biotechnology Co., Ltd. CE 0197 MDD 93/42/EEC Jiangsu HanHeng Medical Technology Co. Ltd. CE 0197 MDD 93/42/FEC Jiangsu Changfeng Medical Industry Co.Ltd.





COVID-19 POSITIVE: Two colored lines appear on the membrane. One line appears in the control region (C) and the other line appears in the test region (T). COVID-19 NEGATIVE: Only one colored line appears in the control region (C). No colored line appears in the test region (T). COVID-19 INVALID: Control line fails to appear.

Influenza A POSITIVE: It is positive for Influenza A antigen if two Red lines appear. One red line should be in the control line region (C), and the other one appears in the A test line region. Influenza B POSITIVE: It is positive for Influenza B antigen if two Red lines appear. One red line should be in the control line region (C), and the other one appears in the B test line region. Influenza A and B POSITIVE: It is positive for both the antigens of Influenza A and Influenza B if three red lines appear. One Red line should be in the control line region (C), and another two should appear in A test line region and B test line region.

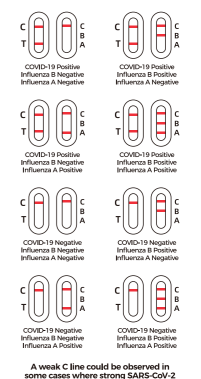
NEGATIVE: One Red line appears in the control region (C). No red line appears in the influenza A and B test region (T). INVALID: Control line fails to appear.

Caution:

- For COVID-19 positive results: If you test positive, follow current Department of Health and Aged Care advice(Refer to FAQ #7).
- For Influenza positive results: If you have a positive result or are unwell, contact a medical practitioner for follow up clinical care.
- For negative results: A negative result does not mean you do not have COVID-19, Influenza A, and/or Influenza B. If symptoms persist or you feel unwell, please contact a medical practitioner for follow up clinical care.
- For Invalid result: Please retest with a new test cassette and a freshly collected specimen. Report repeated invalid results to the sponsor.

The shade of the test line may vary, but even if a faint line appears, this should be considered as positive.

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



positive results are obtained.