



Anti-SARS-CoV-2 Neutralization Antibody Test Kit

(Colloidal gold immunoassay)

Product name: Anti-SARS-CoV-2 Neutralization Antibody Test Kit

(Colloidal gold immunoassay) Catalogue No: CP04005-25 Packing specification: 25T/kit

INTENDED USE

Anti-SARS-CoV-2 Neutralization Antibody Test Kit is a rapid chromatographic immunoassay for semi-quantitative detection of neutralization antibodies to SARS-CoV-2 in human serum or plasma.

SUMMARY

Neutralizing antibody is the most important marker of possible immunity. Anti-SARS-CoV-2 Neutralization Antibody Test Kit is used to detect neutralizing antibodies in serum and plasma, determine whether the body has obtained immunity after vaccine injection, whether the individual that recovered from clinical treatment has potential immunity, and the lifetime of neutralizing antibodies in human body. Coronaviruses encode four major structural proteins, spike (S), membrane (M), envelop (E), and nucleocapsid (N), notably, S protein contains a receptor-binding domain (RBD) which is one of the vital immunodominant epitopes and has a superior capacity to induce neutralizing antibodies. It is proved that RBD of SARS-CoV-2 is responsible for recognizing and interacting with the cell surface receptor, angiotensinconverting enzyme-2 (ACE2). In the respiratory tract, ACE2 is widely expressed on the cell surface of alveoli, trachea, bronchi, macrophages, etc. Following the binding of the RBD to the receptor ACE2, SARS-CoV-2 enters target cells, where the fusion of the virus envelops the endosome membranes and leads to the release of the viral nucleocapsid into the cytosol of the infected cell. Neutralizing antibodies are secreted by B lymphocytes and can bind to virus SP(RBD), which can prevent pathogenic microorganisms from attaching to the receptor ACE2, avoid virus infection.

TEST PRINCIPLE

The Anti-SARS-CoV-2 Neutralizing Antibody Test Kit (Serum/Plasma) is a semi-quantitative membrane-based immunoassay for the detection of SARS-CoV-2 neutralizing antibodies in serum or plasma. the sample is dropped into the sample well, and chromatography is performed under the capillary effect. The SARS-CoV-2 neutralizing antibodies in the sample combined with the colloidal gold-labeled SARS-CoV-2 SP (RBD), then spread to the test area, ACE2 coated on T line is competitively bound to SP (RBD) with neutralizing

antibody in the sample, the more neutralizing antibodies in the sample, the weaker color of T-line showed. The quality control area (C line) is coated with the mouse anti-chicken IgY, and the colloidal gold-labeled chicken IgY is captured to form a complex and aggregate in the C line. If the C line does not show color, it indicates that the result is invalid, and this sample needs to be tested again.

REAGENTS

Test Cassette: 25T/Kit
Buffer: 0.9mL/vial, 25vials

- Desiccant
- Dropper
- Colorimetric Card

STORAGE AND STABILITY

The kit can be stored at 2-30 $^\circ\! \text{C}$ for 18 months. Do not use after the expiration date

SAMPLE COLLECTION AND PREPARATION

Anti-SARS-CoV-2 Neutralization Antibody Test Kit can be performed using serum and plasma.

- Separate serum or plasma from blood as soon as possible to avoid hemolysis. Use only clear, non-hemolyzed samples.
- Test should be performed immediately after sample collection.
- Serum and plasma samples may be stored at 2-8°C for up to 3 days. For long-term storage, samples should be kept below -20°C
- Bring samples to room temperature prior to testing. Frozen samples must be completely thawed and mixed well prior to testing. samples should not be frozen and thawed repeatedly.

PRECAUTIONS

- For medical professional use only.
- Do not reuse.
- Handle all samples cautiously as if they contain infectious agents. Observe established precautions against microbiological hazards throughout the process and properly handle samples in accordance with standard procedures.
- Wear protective clothing such as laboratory coats, disposable gloves and eye protection when samples are assayed.
- The used tests, samples and potentially contaminated should be discarded according to the local regulation.
- Unreasonable sampling, transportation and handling, or low virus content in

the sample will lead to false negative results. Humidity and temperature can adversely affect results.

• Each laboratory should build its own reference range.



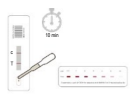
1.Draw 100µL sample.



2.Add 100 µL sample into the buffer.



3.Mix well by vortex.



4.Add 100µL mixture. Compare with the colorimetric card after 10 min.

TEST PROCEDURE

- Bring the pouch to room temperature before opening. Take the cassette from the sealed pouch and use it within ONE hour.
- Place the cassette on a clean and flat surface.
- Add 100 µL of serum or plasma sample into the buffer, mix well, then add 100 µL of mixture and start the timer. Avoid trapping air bubbles into the sample well.
- Wait for the colored line(s) to appear. The result shall be read at 10 minutes.
 The result is valid within 30 minutes.
- Comparing the color shown on T-line with the colorimetric card to get a COI value according to different color grades.

NOTE: It is recommended to use serum or plasma as the priority sample types for testing.

INTERPRETATION OF RESULTS

- If the control line (C line) appears color, please make interpretation with colorimetric card.
- The control line (C line) does not appear color, which means that the test is invalid and the test should be repeated.

^{*} Micropipette is recommended



(6

invalid and the test should be repeated

COI (cut-off index)= 85.299/Value Tine

If result≤1, it means that the signal value of T line is greater than cut-off value, indicating negative.

If result>1, it means that the signal value of T line is less than cut-off value, indicating positive.

PERFORMANCE

- Compliance rate: Result by negative reference COI should be 0.5-1, result by positive reference COI should be 1.5-2.
- 2.Repeatability: ≤15%.
- 3.Specificity analysis: There is no cross-reaction with antibody/antigen positive sera samples from patients with other human coronaviruses (HCoV-HKU1, HCoV-OC43, HCoV-NL63, HCoV-229E), or non-coronaviruses, including influenza A virus (H1N1, H3N2, H5N1, H7N9), influenza B virus (yamagata lineages, victoria lineages), respiratory syncytial virus, rhinovirus, adenovirus, enterovirus, epstein-barr virus, measles virus, human cytomegalovirus, rotavirus, norovirus, mumps virus, herpes zoster virus, or mycoplasma pneumoniae.
- 4.Cut-off: The serum/plasma of 464 healthy people is statistically analyzed, The Cut-off value of "T line" is 85.2998 (Non-parametric percentile method (CLSI C28-A3)).
- Hook effect: There is no hook effect within the titer range of positive samples.
- Clinical performance: 626 clinical samples (111 positive cases and 515 negative cases) showed the sensitivity is 98.198% (95%CI: 93.643%, 99.781%) and specificity is 98.252% (95%CI: 96.709%, 99.198%).

CP04005-25	Plaque-reduction neutralization test		Total	
	Positive	Negative	iolai	
Positive	109	9	118	
Negative	gative 2		508	
Total	111	515	626	
Sensitivity		98.198% (93.643%,99.781%)		
Specificity		98.252% (96.709%,99.198%)		
Positive Predictive Value (PPV)		92.373% (86.366%,95.860%)		
Negative Predictive Value (NPV)		99.606% (98.463%,99.900%)		

Statistics of 111 positive samples (compared with plaque-reduction neutralization test)

Titer	POS By	POS By	Coincidence	CP04005-25 COI		
(PNT50) PNT	PNT	UNscience	rate	Min	Max	Median
<4	14	12	85.71%	0.8	1.3	1.2
4-24	32	32	100%	1.2	1.7	1.4
24-48	22	22	100%	1.4	4.0	2.8
48-96	21	21	100%	2.3	9.9	6.9
>96	22	22	100%	4.9	11.5	8.5

Diagnostic analysis of neutralizing antibodies in different populations:

Characteristic	Total	POS By PNT	POS By UNscience	CP04005-25 COI		
				Min	Max	Median
Heathy	490	0	9	0.6	1.4	0.7
Convalescent	15	6	6	0.7	1.4	0.9
Infestor	21	5	4	0.7	1.5	0.9
Vaccination	100	100	99	0.9	11.5	2.9

LIMITATIONS

- FOR PROFESSIONAL USE ONLY. The presence of binding antibodies does not guarantee the presence of neutralizing antibodies, and the levels can not be consistent.
- 2. The half-life of binding antibody and neutralizing antibody is different, and the decline rate may be different.
- Anti-SARS-CoV-2 Neutralization Antibody Test Kit is for in vitro diagnostic use only. The test should be performed using serum or plasma samples only.
- 4. Anti-SARS-CoV-2 Neutralization Antibody Test Kit will only indicate the presence of SARS-CoV-2 neutralization antibodies in the sample and should not be used as the sole basis, and should be combined with other test methods, as live virus and pseudovirus neutralizing antibodies assay.
- 5. As with all diagnostic tests, all results must be interpreted together with other clinical information available to the physician. Each laboratory should build its own reference range.

BASIC INFORMATION

GLOSSARY OF SYMBOLS

Symbol	Meaning	Symbol	Meaning
IVD	In vitro diagnostic medical device	2℃ - 30℃	Temperature limitation
***	Manufacturer	EC REP	Authorized representative in the European Community
س	Date of Manufacture		Use by date
(2)	Do not reuse	[]i	Consult instructions for use
LOT	Batch code	(6	Meet the requirements of EC Directive 98/79/ EC



Wuhan UNscience Biotechnology Co., Ltd.

Address: Building B18, 2nd Phase of Biomedical Park, #858 GaoXin Road,

Donghu Hi-Tech Development, Wuhan, Hubei, P.R. China Tel: 86-27-87385095

= mail: support@upssions

E-mail: support@unscience.cn



CMC Medical Devices & Drugs S.L

Address: C/Horacio Lengo Nº 18 CP 29006, Málaga-Spain

Tel: +34951214054 Fax: +34952330100

Email: info@cmcmedicaldevices.com Version: 1.0 Date Adopted: 2020-12-01