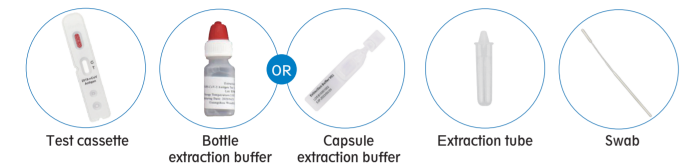
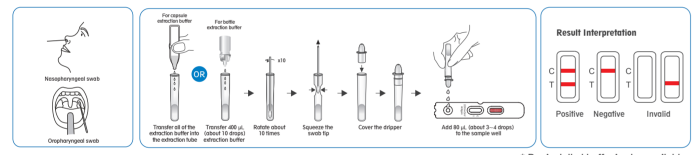
PRODUCT SPECIFICATIONS

Product Components



Operation procedure



* Pre-installed buffer is also available. * The detail of this component (e.g. picture, operation procedure) will be provided upon request.

Performance

Reagents		PCR		Total
Keuyenis		Positive	Negative	Total
Wondfo 2019-nCoV	Positive	208	1	209
Antigen Test (Lateral Flow Method)	Negative	4	361	365
Total		212	362	574

Sensitivity: 98.11% (95%CI: 95.24%~99.48%) Specificity: 99.72% (95%CI: 98.47%~99.99%) Total agreement: 99.13% (95%CI: 97.98%~99.72%)

Order information

Catalog No.	Product Name	Packing Size	Sample Type	Storage Condition	Shelf Life	Qualification
W196	2019-nCoV Antigen Test (Lateral Flow Method)	1T/5T/20T	Nasopharyngeal swab or oropharyngeal swab	2~30 ℃	12 months	CE

Wondfo

Guangzhou Wondfo Biotech Co., Ltd.

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Wondfo

WONDFO BIOTECH WeAreWorkingForYour Heath

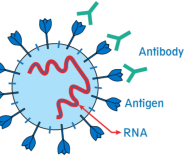
WONDFO 2019-nCoV ANTIGEN TEST

Speed Up the **COVID-19** Control !



WONDFO 2019-nCoV **ANTIGEN TEST**





Antigen test

RT-PCR

Antibody test Detect the antibody generated by immune response after viral infection, indicating the active or past viral infection.

2019-nCo\

WHEN TO USE ANTIGEN TEST?

Releasing profile

*For illustrative purpose only Virus (PCR, Antigen

28 21 35 First infection Symptom onse Antigen

* Incubation Period: 1~14 days, mostly 3~7 days * Antibody Window Period: 5-10 days after onset of symptoms

ANTIGEN TEST ADVANTAGES

Antigen test **OVER** RT-PCR

- Short turn-around time (Antigen test: 20mins vs. RT-PCR: 2hours)
- · Inexpensive cost, no equipment required and simple operation make antigen test suitable for point-of-care (POC) setting usage.

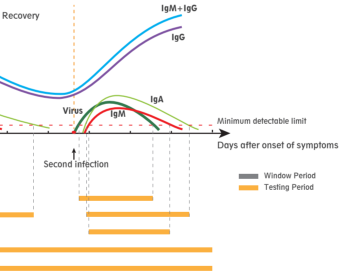
Non-invasive sampling

CURRENT DIAGNOSTIC METHODS FOR COVID-19

Detect the antigen of the virus, indicating the active viral infection.

Detect the RNA of virus, indicating the active viral infection.





* The minimum detectable limit varies with methodology and sensitivity of test * IgG antibody test can be referred as one of discharge criteria for recovering COVID-19 patients

Antigen test **OVER** Antibody test

 Detect the virus directly, allowing the early detection of COVID-19

(sampling type: blood vs. swab)



ANTIGEN TEST APPLICATION

Similar to RT-PCR, the detection of antigen indicates the active infection. Under the circumstance that the area(s) is still undergoing widespread community transmission and/or the area with limited RT-PCR resources, antiaen can be used for aiding in the diagnosis of COVID-19 suspect patients.



* American CDC also recommends to use rapid antiaen tests for screening testing in high-risk congregate settings where the immediate result is required.

Result interpretation



POSITIVE The patient is undergo active 2019-nCoV infection. Further isolation is required.

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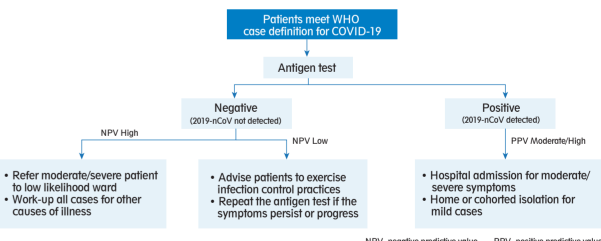
NEGATIVE

The patient should be further evaluated by RT-PCR, especially if the result of the antigen test is inconsistent with the clinical context.

ANTIGEN TEST OFFICIAL GUIDELINES



Antigen-detection in the diagnosis of novel coronavirus (2019-nCoV) infection using rapid immunoassays



NPV- negative predictive value PPV- positive predictive value *The value for NPV and PPV is decided based on products preformance and disease prevalence in applied scenario

Other antigen test related guidelines

- Interim Guidance for Rapid Antigen Testing for 2019-nCoV, American CDC (8-16-20)
- Considerations for Use of 2019-nCoV Antigen Testing in Nursing Homes, American CDC (8-27-20)
- Antigen-Detection in the Diagnosis of 2019-nCoV Infection Using Rapid Immunoassays, WHO (9-11-20)
- Considerations for Implementation of 2019-nCoV Rapid Antigen Testing, APHL (9-2-20)