## Wondfo®

# WONDFO SARS-COV-2 ANTIGEN

TEST

Speed Up the COVID-19 Control!

## **PRODUCT SPECIFICATIONS**

### **Product Components**



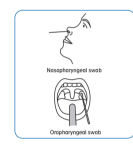


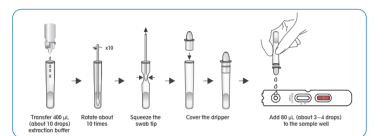


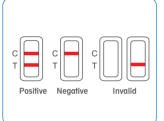




**Operation procedure** 







#### Performance

Reagents		PCR		Takul
Reugeilis		Positive	Negative	Total
Wondfo SARS-CoV-2 Antigen Test (Lateral Flow Method)	Positive	478	1	479
	Negative	19	361	380
Total		497	362	859

Sensitivity: 96.18% (95%CI: 96.43%~98.49%) Specificity: 99.72% (95%CI: 98.45%~99.95%) Total agreement: 97.67% (95%CI: 94.11%~97.54%)

#### **Order information**

Catalog No.	Product Name	Packing Size	Sample Type	Storage Condition	Shelf Life	Qualification
W196	SARS-CoV-2 Antigen Test (Lateral Flow Method)	20T	Nasopharyngeal swab or oropharyngeal swab	2~30 ℃	12 months	C€

Wondfo BIOTECH
WeAreWorkingForYour Health



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# WONDFO **SARS-COV-2 ANTIGEN TEST**





Direct detection of the virus



Room temperature storage(2~30°C)



Instant results within 15mins



Non-invasive sampling (sample type: nasopharyngeal or oropharyngeal swab)



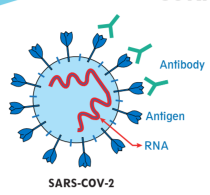


Early detection of COVID-19 (WHO recommends the testing period is from 3 days before to 5-7 days after symptoms onset)

Easy to use, no

equipment required

#### **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.

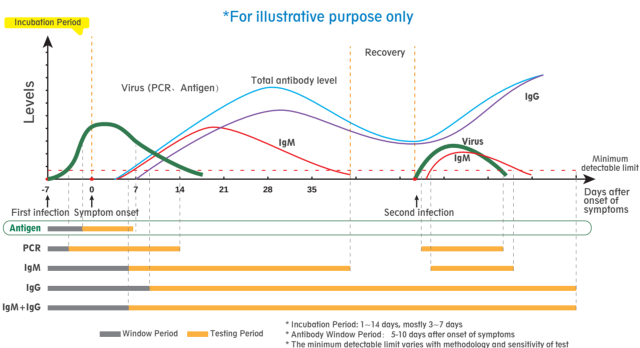
#### **Antibody test**

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

#### WHEN TO USE ANTIGEN TEST?

#### Releasing profile

Levels of SARS-CoV-2 virus and antibodies after infection



#### **ANTIGEN TEST ADVANTAGES**

#### Antigen test **OVER** RT-PCR

- Short turn-ground 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.

#### Antigen test **OVER** Antibody test

- · Detect the virus directly, allowing the early detection of COVID-19
- Non-invasive sampling (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) still undergo widespread community transmission with limited RT-PCR resources, antigen can be used for giding 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 SARS-CoV-2 infection. Further isolation is required.



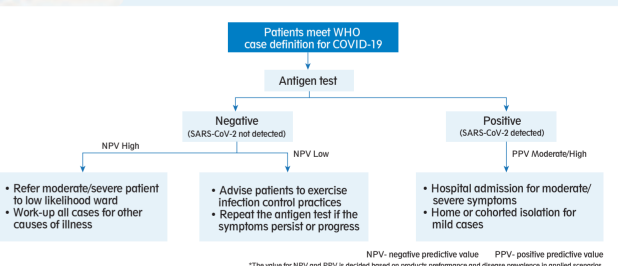
#### NEGATIVE

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 SARS-CoV-2 infection using rapid immunoassays



#### Other antigen test related guidelines

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