

## Rapid Test Kit with 98% Accuracy CE and ISO Certified that use for testing Strep A

Our Product Introduction

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### Basic Information

- Place of Origin: china
- Brand Name: DVOT
- Certification: CE/ISO
- Model Number: Strep A Rapid Test Kit
- Minimum Order Quantity: 800test
- Price: 0.9-1 USD
- Packaging Details: 800T/PER CTN Weight :16.6 KGS CTN SIZE :57\*54\*34CM
- Delivery Time: 15-20 work days
- Payment Terms: D/P, T/T, Western Union, MoneyGram
- Supply Ability: 1000000pcs

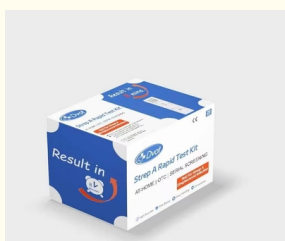


### Product Specification

- Certifications: CE ISO
- Keyword: Antigen Rapid Test
- Result Time: 8 Minutes To Get
- Test Time: 8 Minutes
- Time Of Use: Disposable
- Type: Rapid Test Kit
- Accuracy: 98%
- Highlight: ce clungene antigen rapid test, ce medical diagnostic test kits, disposable clungene antigen rapid test



### More Images



## Product Description

### Rapid Test Kit with 98% Accuracy CE and ISO Certified that use for testing Strep A

#### Product Description:

The Antigen Rapid Self Test Kit is perfect for those who prefer to get tested in the comfort of their own homes, without the need for a medical professional. With this Antigen Home Test Kit, you can easily and quickly test yourself for the presence of viral antigens, helping you to take the necessary precautions and seek further medical advice if needed.

This Antigen Rapid Self Test Kit boasts an accuracy rate of 98%, giving you peace of mind that you are getting reliable and accurate results. With this level of accuracy, you can trust the results you get from the Antigen Rapid Self Test Kit and take the necessary steps to protect yourself and those around you.

With a result time of just 8 minutes, the Antigen Rapid Self Test Kit is one of the fastest diagnostic tools available on the market today. The quick result time allows you to take immediate action and follow the necessary protocols in case of a positive result.

The Antigen Rapid Self Test Kit is designed to be user-friendly and easy to operate. With a test time of just 8 minutes, you can get the results you need in no time. The kit comes with all the necessary components to perform the test, including the test device, swabs, and a buffer solution.

Overall, the Antigen Rapid Self Test Kit is an essential tool for those who want to stay safe and take proactive measures against the spread of viral infections. With its high accuracy rate, quick result time, and user-friendly design, this Antigen Rapid Self Test Kit is a reliable and convenient option for anyone looking to get tested for viral antigens.

#### Technical Parameters:

Test Time	8 Minutes
Accuracy	98%
Time Of Use	Disposable
Certifications	CE, ISO
Type	Rapid Test Kit
Keyword	Antigen Rapid Test
Result Time	8 Minutes To Get

#### Applications:

One of the main advantages of using the DOVT Strep A Rapid Test Kit is that it provides fast and reliable results. The test can be completed in just 8 minutes, which means that healthcare professionals can make an accurate diagnosis and start treatment quickly. This is especially important in cases where patients are showing symptoms of a Strep A infection, such as sore throat, fever, and swollen lymph nodes.

The DOVT Strep A Rapid Test Kit is also very easy to use. It is a disposable test, which means that it can be used once and then discarded. This makes it a very convenient option for healthcare professionals who need to test multiple patients in a short amount of time. The test requires only a small sample of saliva or throat swab from the patient, which is then mixed with the test reagent and inserted into the test cassette. The results are displayed in a clear and easy-to-read format, with two lines indicating a positive result and one line indicating a negative result.

The DOVT Strep A Rapid Test Kit can be used in a variety of different medical settings, including hospitals, clinics, and doctors' offices. It is particularly useful in emergency rooms and urgent care centers where patients need to be diagnosed quickly and accurately. The test is also commonly used in schools and other institutions where there is a risk of Strep A outbreaks.

In summary, the DOVT Strep A Rapid Test Kit is a highly effective and reliable tool for diagnosing Strep A infections. It is a fast, easy-to-use, and disposable test that provides accurate results in just 8 minutes. With its CE and ISO certifications, healthcare professionals can be confident in its quality and reliability. This Antibiotic Test Kit is a must-have for any medical facility that deals with infectious diseases and needs to diagnose patients quickly and accurately.

#### Support and Services:

Our Medical Diagnostic Test Kits are designed to provide accurate and reliable results for a variety of medical conditions. Our technical support team is available to assist with any questions or issues that may arise during the

testing process.

At our company, we are committed to providing the highest level of support and service to our customers. Contact us today to learn more about our Medical Diagnostic Test Kits and the services we offer.

## Advantages:

Conventional identification procedures for Group A Streptococcus from throat swabs involve the isolation and subsequent identification of viable pathogens by techniques that require 24 to 48 hours or longer. DVOT Strep A detects either viable or nonviable organisms directly from a throat swab, providing results within 8 minutes.

DVOT Strep A Rapid Test Kit offers a simple and highly sensitive screening assay to make a presumptive diagnosis of Group A Streptococcal respiratory infection.



## Pathogen and product description

Group A Streptococcal (Strep) infections are caused by Group A Streptococcus, a bacterium responsible for a variety of health problems. These infections can range from mild skin infection or sore throat to severe, life-threatening conditions such as toxic shock syndrome (multi-organ failures) and necrotizing fasciitis (soft tissue disease), commonly known as flesh eating disease. Most people are familiar with strep throat, which along with minor skin infection, is the most common form of the disease. Health experts estimate that more than 10 million mild infections (throat and skin) like these occur every year.

Conventional identification procedures for Group A Streptococcus from throat swabs involve the isolation and subsequent identification of viable pathogens by techniques that require 24 to 48 hours or longer. DVOT Strep A detects either viable or nonviable organisms directly from a throat swab, providing results within 8 minutes.

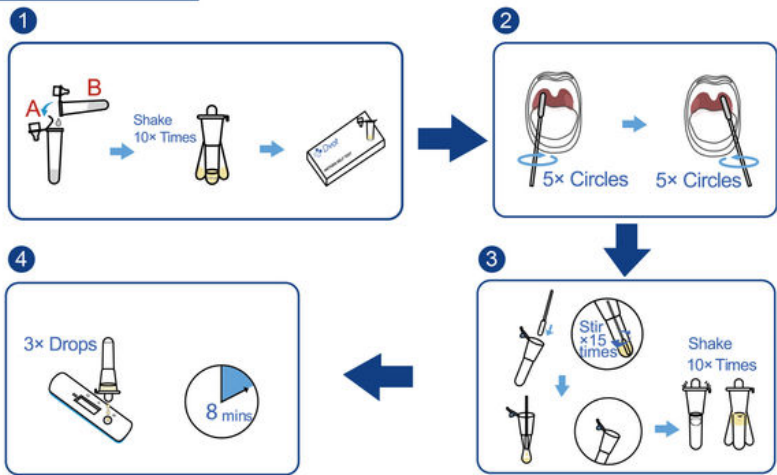
**DVOT Strep A Rapid Test Kit** offers a simple and highly sensitive screening assay to make a presumptive diagnosis of Group A Streptococcal respiratory infection.



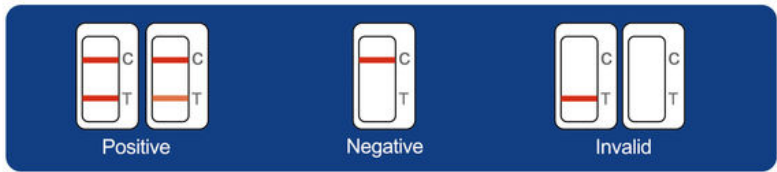
EXCELLENT CLINICAL COMPARISON RESULT

Items	Sensitivity	Specificity
Strep A Rapid Test Kit	92.5%	98.11%

How to test?



Result Reading



The packing details

	
CB	
QTY :	800PCS(40Boxes)
G.W :	15.5KGS
MEAS :	57x54x36cm

# NOTIFICATION OF REGISTRATION

Spanish Medicines and Health Products Administration

This is to certify that, according to the European Council Directive 98/79/EC,  
The manufacturer has completed the registration of the listed products:

**MANUFACTURER:** Feng Chun Yuan Medical Equipment (Shenzhen) Co., Ltd  
**ADDRESS:** Room 1304&1306, No.48, Xinyu Road, Xiangshan Community, Xinqiao  
Street, Baoan District, Shenzhen, China 518000  
**SRN#:** CN-MF-000002326  
**Mail:** market@fcy-medical.com

The manufacturer has provided EU authorized representative with all the  
appropriate declaration according to the European Council Directive 98/79/EC  
including the Declaration of Conformity confirming that its in vitro diagnostic  
medical device, as stipulated here below, is fulfilling the essential requirements of  
the European Council Directive 98/79/EC.

**IVD Devices:** Strep A Rapid Test Kit  
**Classification:** Others

Where the manufacturer affix the CE mark to the device listed they must ensure  
that all the essential requirements of European Council Directive 98/79/EC are met.  
The notification of abovementioned device has been completed by the European  
Authorized Representative in Spain. The Spain Competent Authority is notified of  
the manufacture's device and has allocated registration. The registration number is  
RPS/1389/2024 .



Issue date: 23/MAY/2024  
Cert. No.: RPS/1389/2024

Riomavix S.L. (ES-AR-000001202)  
Calle de Almansa 55, 1D, Madrid 28039 Spain  
Email: info@jiameiconsulting.com

N/REF: PS/RPS/1389/2024

O F I C I O

**Comunicación:** RPS/1389/2024  
**Nº AEMPS:** 24-02001  
**Fecha:** 23/05/2024  
**Asunto:** Anotación de la comunicación  
en el Registro de Responsables  
de la puesta en el mercado de  
Productos Sanitarios

RIOMAVIX SL  
Calle de Almansa 55,1D  
28039 - Madrid  
MADRID  
Madrid, Comunidad de

Con fecha **23/05/2024** ha sido **registrada** en la aplicación de Registro de Responsables de la puesta de mercado de Productos Sanitarios (RPS) de la Agencia Española de Medicamentos y Productos Sanitarios (AEMPS) la comunicación presentada por **RIOMAVIX SL**, con la siguiente información:

1. Número de identificación asignado en el registro

RPS/1389/2024

2. Responsable de la puesta en el mercado de los productos sanitarios

**Empresa** RIOMAVIX SL  
Calle de Almansa 55,1D  
28039 - Madrid (MADRID)  
Madrid, Comunidad de  
**En calidad de** Representante

3. Legislación que declara cumplir:

DIV - Directiva 98/79/EC.

4. Página(s) adicional(es) de productos sanitarios incluidos en esta comunicación.

REGISTRO DE RESPONSABLES DE LA PUESTA EN EL MERCADO DE PRODUCTOS SANITARIOS  
DEPARTAMENTO DE PRODUCTOS SANITARIOS

*Nota.- Esta notificación no tiene el carácter de una autorización sanitaria de comercialización, ni entraña un juicio sobre la conformidad del producto con la legislación vigente. Únicamente avala el cumplimiento del Registro de Responsables según el artículo 9 del RD 1662/2000 por el que se regulan los Productos Sanitarios para Diagnóstico in vitro.*

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 23/05/2024

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: Q3WKLE99AD



Página 1 de 2

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