

## Mycoplasma Pneumoniae IgG/IgM Rapid Test Kit (Colloidal Gold) Instruction for Use

Read this instruction carefully before use

A rapid test for the qualitative detection of Mycoplasma Pneumoniae IgG/IgM in human serum, plasma or whole blood specimens. For professional medical institutions use only, Not for self testing.

#### PRODUCT NAME

Mycoplasma Pneumoniae IgG/IgM Rapid Test Kit (Colloidal Gold)

#### SPECIFICATION

25 tests/kit. 5 tests/kit. 1 test/kit

#### INTENDED USE

The Mycoplasma Pneumoniae Combo Rapid Test is a lateral flow immunoassay for the simultaneous detection and differentiation of IaG and IaM antibody to Mycoplasma Pneumoniae in human serum, plasma or whole blood. It is intended to be used as a screening test and as an aid in the diagnosis of infection COMPONENTS with L. interrogans. Any reactive specimen with the Mycoplasma Pneumoniae IgG/IgM Materials Provided Combo Rapid Test must be confirmed with alternative testing method(s).

#### INTRODUCTION

M.pneumoniae can cause a host of symptoms such as primary atypical pneumonia, tracheobronchitis, and upper respiratory tract disease. Tracheobronchitis is most common in children with a reduced immune system, and up to 18% of infected children require hospitalization, Clinically, M. pneumoniae cannot be differentiated from pneumonia caused by other bacteria or viruses. A specific diagnosis is important because treatment of M. pneumoniae infection with β-lactam antibiotics is ineffective, whereas treatment with macrolides or tetracyclines can reduce the duration of the

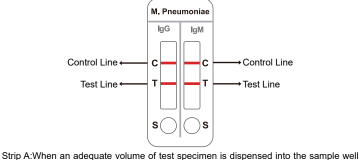
Adherence of M. pneumoniae to the respiratory epithelium is the first step in the infection process. This attachment process is a complex event that requires several adhesin proteins, such as P1, P30, and P116. The true incidence of M. pneumoniae associated infection is not clear as it difficult to diagnose in the early stages of infection.

#### **PRINCIPLE**

Mycoplasma Pneumoniae IgG/IgM Rapid Test Kit based on the principle of a qualitative immunochromatographic assay for the determination of Mycoplasma PRECAUTIONS Pneumoniae IgG/IgM antibody in the in human serum, plasma or whole blood.

StripA consists of: 1) a burgundy colored conjugate pad containing MP antigen • Do not spill solution into the reaction zone. conjugated with colloid gold (MP Antigen conjugates), 2) a nitrocellulose membrane • Do not use test if pouch is damaged. strip containing a test band (T band) and a control band (C band). The T band is pre- • Do not use test kit after expiration date. coated with mouse anti-human IgG antibody, and the C band is pre-coated with goat • Do not mix Sample Diluent Solution and Transfer Tubes from different lots. anti-mouse IaG antibody.

Strip B consists of : 1) a burgundy colored conjugate pad containing MP antigen • Do not spill solution into the reaction zone. conjugated with colloid gold (MP Antigen conjugates), 2) a nitrocellulose membrane • For professional use only. strip containing a test band (T band) and a control band (C band). The T band is precoated with mouse anti-human IaM antibody, and the C band is pre-coated with goat • Do not touch the reaction zone of the device to avoid contamination. anti-mouse IgG antibody.



of the test cassette, the specimen migrates by capillary action across the cassette. MP IgG antibody if present in the specimen will bind to the MP Antigen conjugates. The immunocomplex is then captured on the membrane by the pre-coated Mouse anti-human IgG antibody, forming a burgundy colored T band, indicating a MP IgG positive test result. Absence of the T band suggests a negative result. The test contains an internal control (C band) which should exhibit a burgundy colored band of the immunocomplex of goat anti-mouse IgG/mouse IgG-gold conjugate regardless of the presence of colored T band. Otherwise, the test result is invalid and the specimen must be retested with another device.

Strip B:When an adequate volume of test specimen is dispensed into the sample well of the test cassette, the specimen migrates by capillary action across the cassette. MP IgM antibody if present in the specimen will bind to the MP Antigen conjugates. The immunocomplex is then captured on the membrane by the pre-coated Mouse anti-human IgM antibody, forming a burgundy colored T band, indicating a MP IgM positive test result. Absence of the T band suggests a negative result. The test contains an internal control (C band) which should exhibit a burgundy colored band of the immunocomplex of goat anti-mouse IgG/mouse IgG-gold conjugate regardless of the presence of colored T band. Otherwise, the test result is invalid and the specimen must be retested with another device.

Components	25 tests/kit	5 tests/kit	1 tests/kit
Cassettes	25 cassettes with dependent sealed foil pouch	5 cassettes with dependent sealed foil pouch	1 cassette with dependent sealed foil pouch
Sample Diluent Solution	Sample Diluent Solution 25tubes(300ul/ tube) 5tubes(300ul/ tube)		300ul/tube
Dropper	25 pcs	5 pcs	1 pcs
Lancet	25 pcs	5 pcs	1 pcs
Alcohol pad	25 pcs	5 pcs	1 pcs
Package insert 1		1	1

Main ingredients of Sample Diluent Solution:

Neutral salt buffer

Reagents of different batch numbers cannot be used interchangeably.

#### MATERIALS REQUIRED BUT NOT PROVIDED

Timer for timing use

- · Read this IFU carefully before use.

- Do not open the Test Cassette foil pouch until ready to perform the test.

- · For in-vitro diagnostic use only
- Avoid cross-contamination of samples by using a new specimen collection container and specimen collection tube for each sample.
- · All patient samples should be treated as if capable of transmitting disease. Observe established precautions against microbiological hazards throughout testing and follow standard procedures for proper disposal of specimens.
- Do not use more than the required amount of liquid.
- Bring all reagents to room temperature (15~30°C) before use.
- · Wear protective clothing such as laboratory coats, disposable gloves and eye protection when testing.
- Evaluate the test result after 20 minutes and not beyond 30 minutes.
- Store and transport the test device always at 2~30°C.

#### STORAGE AND STABILITY

- The kit should be stored at 2~30°C, valid for 12months.
- The test must remain in the sealed pouch until use
- · Do not freeze.

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- Cares should be taken to protect components in this kit from contamination
- Do not use if there is evidence of microbial contamination or precipitation. Biological contamination of dispensing equipment, containers or reagents can lead to false

#### ASSAY PROCEDURE

- Step 1: Bring the specimen and test components to room temperature if refrigerated or frozen. Once thawed, mix the specimen well prior to assay.
- Step 2: When ready to test, open the pouch at the notch and remove device. Place the test device on a clean, flat surface.
- Step 3: Be sure to label the device with specimen's ID number.

### Step 4: For whole blood test

- Apply 1 drop of whole blood (about 30 µL) into the sample well.
- Then add 2 drops (about 60-70 µL) of Sample Diluent immediately.



#### For serum or plasma test

- Fill the pipette dropper with the specimen.
- Holding the dropper vertically, dispense 1 drop (about 30-35 uL) of specimen into the sample well making sure that there are no air bubbles.
  - Then add 2 drops (about 60).



Step 5: Set up timer.

Step 6: Results can be read in 20 minutes. Positive results can be visible in as short as 1 minute.

Don't read results after 30 minutes

#### QUALITY CONTROL

Internal Control: This test contains a built-in control feature, the C band. The C line develops after adding specimen and sample diluent. Otherwise, review the whole procedure and repeat test with a new device.

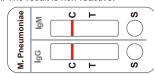
External Control: Good Laboratory Practice recommends using the external controls, positive and negative, to assure the proper performing of the assay, in particularly, under the following circumstances:

- a. New operator uses the kit, prior to performing testing of specimens.
- b.A new lot of test kit is used.
- c.A new shipment of kits is used.
- d. The temperature used during storage of the kit fall outside of 2~30°C.
- e.The temperature of the test area falls outside of 15~30°C.

#### INTERPRETATION OF ASSAY RESULT

#### **Negative Control**

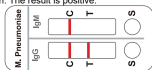
If only the C band is developed, the test indicates that no detectable MP IgG/IgM is present in the specimen. The result is non-reactive.





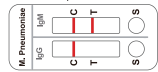
# Positive Control

If both C and IgG lines are developed, the test indicates the presence of MP IgG antibody in the specimen. The result is positive.



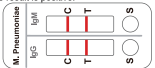
#### MP IqM positive

If both C and IgM lines are developed, the test indicates the presence of MP IgM antibody in the specimen. The result is positive.



#### IgG and IgM antibody positive

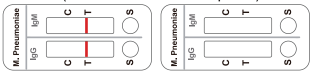
If both C, IgG and IgM lines are developed, the test indicates the presence of MP IgG/IgM in the specimen. The result is positive.



#### INVALID:

If no C line is developed, the assay is invalid regardless of any color development on the T line as indicated below. Repeat the assay with a new test device.

Excess fecal specimen can lead to invalid test results; if this is the cause, resample and re-test (see instructions for collection of specimen).



The appearance of any burgundy color in the test bands, regardless of intensity, must be considered as presence of the band.

Samples with positive or reactive results should be confirmed with alternative testing method(s) such as ELISA or PCR and clinical findings before a diagnostic decision is made

#### PERFORMANCE CHARACTERISTICS

#### 1. Clinical Performance For IgM Test

A total of 210 samples from susceptible subjects were tested by the Mycoplasma Pneumoniae IgG/IgM Combo Rapid Test and by a commercial Mycoplasma Pneumoniae IgM EIA kit. Comparison for all subjects is showed in the following table.

	Mycoplasma Pne Combo R			
IgM EIA	Positive	Negative	Total	
Positive	9	1	10	
Negative	2	198	200	
Total	11	199	210	
Relative Sensitivity: 90.0%, Relative Specificity: 99.0%,				

Overall Agreement: 98.6%

#### 2. Clinical Performance For IgG Test

A total of 206 samples from susceptible subjects were tested by the Mycoplasma Pneumoniae IgG/IgM Combo Rapid Test and by a commercial Mycoplasma Pneumoniae IgG EIA kit. Comparison for all subjects is showed in the following table.

	Mycoplasma Pne Combo R			
IgG EIA	Positive	Negative	Total	
Positive	6	0	6	
Negative	2	2 198		
Total	8	198	206	
Relative Sensitivity: 100%, Relative Specificity: 99.0%,				

#### **TEST LIMITATIONS**

1.The Assay Procedure and the Test Result Interpretation must be followed closely when testing the presence of antibodies to pathogenic Mycoplasma Pneumoniae in serum, plasma or whole blood from individual subjects. Failure to follow the procedure may give inaccurate results.

Overall Agreement: 99.0%

- 2.The Mycoplasma Pneumoniae IgG/IgM Rapid Test is limited to the qualitative detection of antibodies to Mycoplasma Pneumoniae in human serum, plasma or whole blood. The intensity of the test band does not have linear correlation with antibody titer in the specimen.
- 3.A negative result for an individual subject indicates absence of detectable Mycoplasma Pneumoniae antibodies. However, a negative test result does not preclude the possibility of exposure to Mycoplasma Pneumoniae.
- 4.A negative result can occur if the quantity of Mycoplasma Pneumoniae antibodies present in the specimen is below the detection limits of the assay, or the antibodies that are detected are not present during the stage of disease in which a sample is collected
- 5. Some specimens containing unusually high titer of heterophile antibodies or rheumatoid factor may affect expected results.
- 6.The results obtained with this test should only be interpreted in conjunction with other diagnostic procedures and clinical findings.

### SYMBOLS

Symbol	Used For	Symbol	Used For		
	Use-by date	(i	Consult instructions for use		
LOT	Batch code	IVD	In vitro diagnostic medical device		
1	Temperature limit		Manufacturer		
2	Please don't reuse it	*	Keep away from sunlight		
<b>®</b>	Don't use the product when the package is damaged	<del>*</del>	Keep dry		
	Date of manufacture	Σ	Tests per kit		
CE	CE Mark	\$	Biological Risks		
EC REP	Authorized representative in the European Community				

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#### BASIC INFORMATION



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