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Giardia Antigen Rapid Test Kit

(Fecal Specimen)

Instruction for Use

Read this instruction carefully before use

A rapid test for the qualitative detection of Human Giardia antigen in human fecal specimens. For professional medical institutions use only. Not for self testing.

PRODUCT NAME

Giardia Antigen Rapid Test Kit (Fecal Specimen)

SPECIFICATION

40 tests/kit, 25 tests/kit, 5 tests/kit

INTENDED USE

The Giardia Antigen Rapid Test Kit is a lateral flow chromatographic immunoassay for the qualitative detection of Giardia in human Fecal Specimen. It is suitable for the Main ingredients of test cassettes: auxiliary diagnosis of Giardia infection.

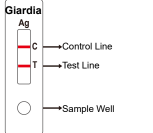
INTRODUCTION

Giardia, a flagellated protozoan, inhabits the upper part of the small intestine of its host and has a two major states in the life cycle:trophozoites which produces the antigens(α -1 giardin) and cyst with produces the antigens(CWP1). After the host ingest the cysts, which are the infective stage, the trophozoites emerge from the cysts in the duodenum and attach to the small intestinal mucosa. They undergo mitotic division in the intracellular lumen, some will encyst to protect themselves and will be eliminated from the host in the feces. The trophozoite is the vegetative form and replicates in the small intestine.

Giardiasis is a diarrheal illness caused by a very small parasite Giardia intestinalis(also • Do not spill solution into the reaction zone. known as Giardia lamblia and Giardia duodealis). Once an animal or person is infected • Do not use test if pouch is damaged. with Giardia, the parasite lives in the intestine and is passed in the stool. The parasite • Do not use test kit after expiration date. is protected by an outer shell and can survive outside the body and in the environment • Do not mix Sample Diluent Solution and Transfer Tubes from different lots. for a long time. The most common symptoms of giardiasis include: diarrhea. loose or • Do not open the Test Cassette foil pouch until ready to perform the test. watery stool, stomach cramps and upset stomach. These symptoms generally begin • Do not spill solution into the reaction zone. 1-2 weeks after infection, and may last 2-6 weeks in healthy indiciduals. Sometimes . For professional use only. symptos last longer and may lead to weight loss and dehydration.Some people will have no symptos. However, people with weakened immune systems (e.g. persons with • Do not touch the reaction zone of the device to avoid contamination. HIV/AIDS, camcer patients, and transplant patients) ot the elderly may have a more • Avoid cross-contamination of samples by using a new specimen collection container serious infection that can lead to severe illness or death.

PRINCIPLE

The Giardia Antigen Rapid Test Kit is a lateral flow chromatographic immunoassay. The test cassette consists of: 1) a burgundy colored conjugate pad containing recombinant antigen conjugated with colloid gold (monoclonal mouse anti-Giardia antibody conjugates) and rabbit IgG-gold conjugates, 2) a nitrocellulose membrane strip containing test band (T bands) and a control band (C band). The T band is precoated with monoclonal mouse anti-Giardia antibody for the detection of Giardia



antigen, and the C band is pre-coated with goat anti rabbit IgG. When an adequate

volume of test specimen is dispensed into the sample well of the test cassette, the

Giardia if present in the specimen will bind to the monoclonal mouse anti-Giardia

the pre-coated mouse anti-Giardia antibody, forming a burgundy colored T band,

specimen migrates by capillary action across the cassette.

indicating a Giardia antigen positive test result. Absence of test band (T) 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 rabbit IgG/rabbit IgG-gold conjugate regardless of the color development on any of the test bands. Otherwise, the test result is invalid, and the specimen must be retested with another device.

COMPONENTS

Materials Provided

Components	40 tests/kit	25 tests/kit	5 tests/kit
Cassettes		25 cassettes with dependent sealed foil pouch	5 cassette with dependent sealed foil pouch
Sample Diluent Solution 1mL/bottle, 40pcs		1mL/bottle, 25pcs	1mL/bottle, 5pcs
Transfer tube 40 pcs		25 pcs	1 pcs
Package insert 1 pcs		1 pcs 1 pcs	

Mouse anti-Human Transferrin antibody, Goat anti-rabbit IgG polyclonal antibody, Human Transferrin antibody, rabbit IgG, Colloidal gold conjugate, Other test device support: one desiccant.

Main ingredients of Sample Diluent Solution:

Neutral salt buffer

Reagents of different batch numbers cannot be used interchangeably. MATERIALS REQUIRED BUT NOT PROVIDED

for timing use

PRECAUTIONS

Timer

- Read this IFU carefully before use.

- · For in-vitro diagnostic use only
- 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.

· 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 results.

SPECIMEN COLLECTION AND HANDLING

Consider any materials of human origin as infectious and handle them using standard biosafety procedures.

To prepare specimens using solid stool samples follow Procedure A below. To prepare specimens using watery stool samples follow Procedure B below.

Procedure A: Solid stool samples

antibody conjugates. The immunocomplex is then captured on the membrane by Step 1: Collect a random stool sample in a clean, dry receptacle.

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Step 2: Open the stool collection device by unscrewing the top and use the collection stick to randomly pierce the stool sample in at least five different sites. Do not scoop stool sample as this may lead to an invalid test result.

Step 3: Ensure stool sample is only in the grooves of the collection stick. Excess stool sample may lead to an invalid test result.

Step 4: Replace the collection stick and tighten securely to close the stool collection device.

Step 5: Shake the stool collection device vigorously.



Procedure B: Watery stool samples

Step 1: Collect a random stool sample in a clean, dry receptacle.

Step 2: Open the stool collection device by unscrewing the top.

Step 3: Fill the plastic dropper with the sample; dispense 2 drops (70-85uL) into the stool collection device.

Step 4: Replace the collection stick and tighten securely to close the stool collection device.

Step 5: Shake the stool collection device vigorously.

Note: Specimens extracted may be stored at 2-8°C for up to 3 days. If longer storage is required, freezing at ≤-20°C is recommended.

ASSAY PROCEDURE

Step 1: Bring the specimen and test components to room temperature if refrigerated or frozen.

Step 2: When ready to test, open the pouch at the notch and remove the test device. Place the test device on a clean, flat surface.

Step 3: Shake the stool collection device vigorously to ensure an effective liquid suspension.



Step 4: Position the stool collection device upright and twist off the dispenser cap. Holding the stool collection device vertically, dispense 2 drops of the solution (85-95uL) into the sample well of the test device. Do not overload sample.

Step 5: Set up timer.

Step 6: Results can be read after 15 minutes. Positive results can be visible in as short as 1 minute.

Don't read results after 30 minutes. To avoid confusion, discard the test device

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after interpreting the result.

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°C -30°C.

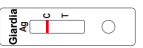
e.The temperature of the test area falls outside of 15°C -30°C.

INTERPRETATION OF ASSAY RESULT

Negative Control

If only the C band is developed, the test indicates that no detectable Giardia antigen is present in the specimen. The result is non-reactive.

Positive Control:

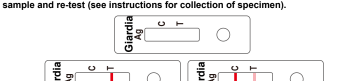


If both C and T lines are developed, the test indicates the presence of Giardia antigen in the specimen. The result is positive.



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, re**-



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.Sensitivity, Specificity and Accuracy

A total of 610 patient samples from susceptible subjects were test by the ELISA test. Comparison for all subjects is showed in the following table:

Giardia Antigen Test	ELISA Test				
BESTest	Positive	Negative	Total		
Positive	108	2	110		
Negative	2	498	500		
Total	110	500	610		
Relative Sensitivity: 98.18%; Relative Sp ecificity:99.6%; Overall agreement:					
99.34%					

2.Cross-reactivity

An evaluation was performed to determine the cross reactivity of BESTest Astrovirus , no cross reactivity against gastrointestinal pathogens occasionally present in faeces:

Name	Name	Name	Name
Staphylococcus aureus	Enterovirus	Listeria monocytogenes	Staphylococcus aureus
Campylobacter coli	Entamoeba hystolitica	Salmonella typhi	Shigella sonnei
Campylobacter jejuni	Escherichia coli O157:H7	Salmonella typhimurium	Shigella dysenteriae
Clostridium Difficlie	Cryptosporidium parvum	Salmonella enteritidis	Shigella flexneri
Shigella sonnei	Helicobacter pylori	Salmonella paratyphi	

3.Interfering Substances

This kit has no interference with HAMA, Human serum Albumin, Antinuclear antibody, Antimitochondrial antibody, Cholesterol, Bilirubin conjugated, Lipids, Hemoglobin, Bilirubin unconjugated, Rheumatoid factor, et al.

QUALITY CONTROL

1.Internal procedural controls are included in the test. A colored band appearing in the control region (C) is considered an internal positive procedural control. It confirms sufficient specimen volume and correct procedural technique.

2.External controls are not supplied with this kit. It is recommended that positive and negative controls be tested as a good laboratory practice to confirm the test procedure and to verify proper test performance.

TEST LIMITATIONS

1. The Giardia Antigen Rapid Test Kit (Fecal Specimen) is for in vitro diagnostic use only. This test should be used for the detection of Giardia antigens in human Fecal specimens.

2. The Giardia Antigen Rapid Test Kit (Fecal Specimen)will only indicate the presence to Cryptosporidium in the specimen and should not be used as the sole criteria for the diagnosis of Giardia infections.

3.If the symptom persists, while the result from Giardia Antigen Rapid Test is negative or non-reactive result, it is recommended to re-sample the patient few hours later.

4.As with all diagnostic tests, all results must be interpreted together with other clinical information available to the physician.

5.If the test result is negative and clinical symptoms persist, additional testing using other clinical methods is recommended. A negative result does not at any time preclude the possibility of Giardia infection.

6.The potential impacts of vaccines, antiviral therapeutics, antibiotics, chemotherapeutic or immunosuppressant drugs have not been evaluated in the test. 7.Due to inherent differences between methodologies, it is highly recommended that, prior to switching from one technology to the next, method correlation studies are undertaken to qualify technology differences. One hundred percent agreement between the results should not be expected due to differences between technologies. 8.Performance has only been established with the specimen types listed in the Intended Use. Other specimen types have not been evaluated and should not be used

with this assay.

CAUTION

1. This product is used for in vitro diagnosis only.

2.Must strictly follow the instructions for operation and interpretation of the results. 3.The product is qualitatively tested, and the result cannot be used as a quantitative basis.should be tested using reagents within the validity period.

4.The cassetes, collectors,droppers,and tubes are for single person one-time use, cannot be reused.

5.Because the sample titer is different, the red lines of the test line will show different shades of color, all of which indicate positive results. The depth of the test line color cannot be used as the basis for determining the antibody titer in the sample.

6.The samples stored at low temperature should be balanced to room temperature and fully mixed before testing.

7.Samples and waste must be treated as a potential source of infection and the desiccant in the foil bag is not edible.

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SYMBOLS				
Symbol	Used For	Symbol	Used For	
	Use-by date	ĺ	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	
	Don't use the product when the package is damaged	Ť	Keep dry	
	Date of manufacture	Σ	Tests per kit	
CE	CE Mark	Ŕ	Biological Risks	
EC REP	Authorized representative in the European Community			

BASIC INFORMATION



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