HI-Quik Typhoid IgG/IgM Rapid Card Test (Serum/Plasma) Package Insert

A rapid test for the qualitative detection of IqG and IqM antibodies to Salmonella typhi (S. typhi) in human serum or plasma.

For professional in vitro diagnostic use only

[INTENDED USE]

The HI-Quik Typhoid IgG/IgM Rapid Card Test is a rapid chromatographic immunoassay for the simultaneous detection and differentiation of IgG and IgM types of antibodies against Salmonella typhi (S. typhi) in human serum or plasma. It is intended to be used as a screening test as an aid in the diagnosis of infection with S typhi. Any reactive specimen with the Typhoid rapid test cassette needs to be confirmed with alternative testing method.

[SUMMARY]

Typhoid fever is caused by S. typhi, a Gram-negative bacterium. World-wide an estimated 17 million cases and 600 000 associated deaths occur annually Patients who are infected with HIV are at significantly increased risk of clinical infection with S typhi². Evidence of h. pylori infection also presents an increase risk of acquiring typhoid fever. 1-5% of patients become chronic carrier harboring S. typhi in the gallbladder

The clinical diagnosis of typhoid fever depends on the isolation of S. typhi from blood, bone marrow or a specific anatomic lesion in the facilities that cannot afford to perform this complicated and time consuming procedure. Widal Test (also referred as Weil-Felix Test) is used to facilitate the diagnosis. However, many limitations lead to difficulties in the interpretation of the Widal test ^{3, 4}.

In contrast, the Typhoid Rapid Test Cassette is a simple and rapid laboratory test. The test simultaneously detects and differentiates the IgG and the IgM antibodies to S. typhi specific antigen⁵ in serum or plasma thus aid in the determination of current or previous exposure the S. typhi.

[PRINCIPLE]

The HI-Quik Typhoid IgG/IgM Rapid Card Test is a qualitative, membrane based immunoassay for the detection of antibodies (IgG and IgM) to Salmonella typhi (S typhi) in human serum or plasma. The diagnostic test cassette consists of two components; an IgG component and an IgM component. The IgG line region is precoated with reagents for the detection of anti-S, typhi (IgG). The IgM line region is precoated with monoclonal anti-human IgM for detection of anti-S, typhi (IgM).

During testing, specimen dispensed into the sample well of the test cassette binds with Typhoid conjugates impregnated in the reagent area, if the specimen contains anti-Typhoid antibodies. The immunocomplex thus formed migrates by capillary action. If the present antibodies in specimen are of IgG types, the immunocomplex is then captured by the pre-coated reagents on the membrane, forming a colored IgG line, indicating a S. typhi IgG positive test result. If the present antibodies in the specimen are of IgM type, the immunocomplex would be captured on the membrane by the pre-coated anti-human IgM antibody, forming a colored IgM line, indicating a S. typhi IgM positive test result.

Absence of any T lines (IgM and IgG) indicates a negative result. A colored control line (C) should always appear in case of a positive or a negative result. Its absence indicates invalid test results.

[REAGENTS]

The test contains mouse anti-human IgM, mouse anti-human IgG and Typhoid antigen. A goat antibody is employed in the control line system.

[PRECAUTIONS]

- 1. For in vitro diagnostic use only. Do not use after the expiration date.
- 2. Do not smoke, drink, or eat in areas where specimens or kits reagents are handled.
- 3. Dispose of all specimens and materials used to perform the test as biohazardous
- 4. This package insert must be read completely before performing the test.
- Bring all reagents to room temperature (15°C-30°C) before use.
- 6. Do not interchange the buffer and test cassettes of different lots.
- 7. Do not use hemolized blood specimen for testing.

[STORAGE AND STABILITY]

Store as packaged in the sealed pouch at room temperature or refrigerated (2-30°C). The test is stable through the expiration date printed on the sealed pouch. The test must remain in the sealed pouch until use. DO NOT FREEZE. Do not use beyond the expiration date

[SPECIMEN COLLECTION AND PREPARATION]

- 1. The HI-Quik Typhoid IgG/IgM Rapid Card Test can be performed using serum or
- 2. Separate the serum or plasma from blood as soon as possible to avoid hemolysis. Only clear, non-hemolyzed specimens can be used.
- 3. Testing should be performed immediately after the specimens have been collected. Do not leave the specimens at room temperature for prolonged periods. Specimens may be stored at 2-8°C for up to 3 days, for long-term storage. specimens should be kept below -20°C.
- 4. Bring specimens to room temperature prior to testing. Frozen specimens must be completely thawed and mixed well prior to testing. Specimens should not be frozen and thawed repeatedly.
- 5. Do not use samples demonstrating gross lipemia, gross hemolysis or turbidity.
- 6. If specimens are to be shipped, they should be packed in compliance with local regulations.

[MATERIALS]

Materials provided • Test cassettes

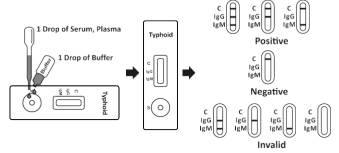
 Sample droppers • Buffer Package insert Materials required but not provided

Specimen collection containers

•Timer • Centrifuge

[DIRECTIONS FOR USE]

- 1. Bring the pouch to room temperature before opening it. Remove the test cassette from the sealed pouch and use it as soon as possible. Best results will be obtained if the assay is performed within one hour.
- 2 Place the test cassette on a clean and level surface. Hold the dropper vertically and transfer 1 drop of serum or plasma (25 ul) to the specimen well of the test cassette, then add 1 drop of buffer (40 ul) and start the timer. Avoid trapping air bubbles in the specimen well. See the illustration below.
- 3. Wait for the colored line(s) to appear. Read the result at **15 minutes**; do not interpret the result after 20 minutes.



[INTERPRETATION OF RESULTS] (Please refer to the illustration above) POSITIVE:* Two or three lines appear. One colored line should always appear in the control line region (C) and another one or two apparent colored line(s) should be in the test line region(s) (IgM and/or IgG).

IgM Positive: Along with line in Control region (C), a line appears in IgM region. It indicates a positive Test result for antibodies to S. typhi (Isotype IgM)

laG Positive: Along with line in Control region (C), a line appears in laG region. It indicates a positive Test result for antibodies to S. typhi (Isotype IgG)

*NOTE: The intensity of the color in the test line regions (IgM and IgG) may vary depending on the concentration of Typhoid antibodies present in the specimen. Therefore, any shade of color in the test line region (IgM and/or IgG) should be considered positive.

NEGATIVE: One colored line appears in the control line region (C). No line appears in the test line regions (IgM and IgG).

INVALID: Control line fails to appear. Insufficient specimen volume or incorrect procedural techniques are the most likely reasons for control line failure. Review the procedure and repeat the test with a new test. If the problem persists, discontinue using the test kit immediately and contact your local distributor.

[QUALITY CONTROL]

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

- 1. The assay procedure and the test result interpretation must be followed closely when performing the assay. Failure to follow the procedure may give inaccurate
- 2. The HI-Quik Typhoid IgG/IgM Rapid Card Test is for qualitative detection of antibodies to S. typhi in human serum or plasma. The intensity of the test band has not linear correlation with the antibody titer in the specimen.
- 3. A negative result only indicates absence of anti-S. typhi antibodies above detectable levels. A negative test result does not preclude the possibility of exposure to S. typhi as a negative result can occur if the quantity of anti-S typhi antibodies present in the specimen is below the detection limit of the assay, or the antibodies that are detected are not present during the stage of disease in which a sample is collected.
- 4. Specimens containing unusually high titer of heterophile antibodies or rheumatoid factor may affect expected results.
- 5. The results obtained with this test should only be interpreted in conjunction with other diagnostic procedures and clinical findings.

[EXPECTED VALUES]

The HI-Quik Typhoid IgG/IgM Rapid Card Test (Serum and Plasma) has been compared with a leading commercial Typhoid ELISA test. The correlation between these two systems is over 96%

[PERFORMANCE CHARACTERISTICS]

Sensitivity and Specificity

A clinical evaluation was conducted comparing the results obtained using the HI-Quik Typhoid IgG/IgM Rapid Card Test to Typhoid IgG/IgM ELISA Testing. The study

included 367 IgG specimens and 230 IgM specimens, and about the IgG specimen both assays identified 333 negative and 25 positive results, about the IgM specimen both assays identified 185 negative and 36 positive results.

laG Results Method S. typhi FLISA (laG) Total Results Results Positive Negative Typhoid Rapid Test 32 Positive 25 Cassette for IgG Negative 333 335 2 340 Total Results 367

Sensitivity: 92.6% (95%CI*: 75.7%~99.1%).Specificity: 97.9% (95%CI*: 95.8%~99.2%) Accuracy: 97.5% (95%CI*: 95.4%~98.9%) *Confidence Intervals

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Method			S. typhi ELISA (IgM)		Total Results		
Typhoid Rapid Test Cassette for IgM	Toot	Results	Positive	Negative	Total Results		
		Positive	36	5	41		
	givi	Negative	4	185	189		
Total Results			40	190	230		

Sensitivity: 90.0% (95%CI*: 76.3%~97.2%), pecificity: 97.4% (95%CI*: 94.0%~99.1%) Accuracy: 96.1% (95%CI*: 92.7%~98.2%) *Confidence Intervals

Precision Intra-Assay

Within-run precision has been determined by using 10 replicates of three specimens: a negative, a low positive, and a high positive. The negative, low positive, and high

positive values were correctly identified >99% of the time. Inter-Assav Between-run precision has been determined by 10 independent assays on the same

three specimens: a negative, a low positive, and a high positive. Three different lots of the Typhoid Rapid Test cassette (Serum/Plasma) have been tested over a 3-day period using negative, low positive, and high positive specimens. The specimens were correctly identified >99% of the time

Cross-reactivity

The HI-Quik Typhoid IgG/IgM Rapid Card Test (Serum/Plasma) has been tested for HBsAg, HBsAb, HbeAg, HBeAb, HBcAb, HCV, HIV, Syphilis, H. Pylori, CMV, Rubella and Toxo positive specimens. The results showed no cross-reactivity.

Interfering Substances

The following potentially interfering substances were added to Typhoid negative and positive specimens.

Caffeine: 20 mg/dL Acetaminophen: 20 mg/dL Acetylsalicylic Acid: 20 mg/dL Gentisic Acid: 20 mg/dL Ascorbic Acid: 2a/dL Albumin: 2 a/dL Bilirubin: 1a/dL Oxalic Acid: 600mg/dL

None of the substances at the concentration tested interfered in the assay.

[BIBLIOGRAPHY]

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[Symbol Legends]

[Oylinbol Legerius]							
S ym bo l	E x pla nat ion o f Sy m bo l	S ymb ol	E x pla inat ion of S y m b ol				
[]i	Co n su It inst ru ct ion fo r use	8	D o n ot re -us e				
8	Do not use if package is damaged	Σn	Contains sufficient for "n" tests				
IVD	In v itro d ia g n o s t ic d e v ic e	REF	Catalognumber				
X	Store at 2 ° c - 30 ° c	LOT	Batchcode				
~~~	D ate of m an uf ac ture	***	Manufacturer				
2<	Use by (date or month of expiry)						

## Manufactured by:

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