

MULTISURE Dengue Ab/Ag Rapid Test

Instructions For Use

FOR RESEARCH USE ONLY NOT FOR USE IN DIAGNOSTICS PROCEDURES

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43592-020 (20 tests)

Note: Changes highlighted

TRADE NAME AND INTENDED USE

The **MP Diagnostics MULTISURE Dengue Ab/Ag Rapid Test** is a qualitative *in vitro* immunochromatographic test to detect and differentiate IgA/ IgM/ IgG antibodies against dengue virus and NS1 dengue antigen in human plasma, serum, finger pricked whole blood or whole blood with anti-coagulants.

This kit is supplied for research purposes only. It is not intended for use in the diagnosis or prognosis of disease.

INTRODUCTION

Dengue fever, a breakbone fever, is a mosquito-borne tropical disease caused by the dengue virus, transmitted by several species of mosquito within the genus *Aedes*, mainly *A.argypti*. There are four different serotypes which all cause clinical disease. Dengue virus is now circulating in Asia, Africa and Americas, a dramatically different scenario from that which prevailed 20 or 30 years ago ^[1, 3]. It is estimated that more than 2.5 billion people live in dengue endemic countries are at risk of infection ^[1, 2].

In general, diagnostic of dengue is dependent on the phase of the infection, current dengue diagnostics are based on either detection of viral agent (antigen/genome) or antibodies (IgA/IgM/IgG) produced against it [3, 9]. Dengue nonstructural protein 1 (NS1), produced in both membraneassociated and secreted forms, has been utilized as an early marker due to its presence in the serum of patients during the viraemic phase of infection. NS1 circulates in the serum from 1 to 9 days after the onset of clinical signs, with a peak from 3 to 5 days [4, 5]. The host immune response to dengue infection consists of the production of antibodies that are primarily directed against the dengue virus envelope proteins ^[3]. Understanding the features of host humoral immune response is important for the interpretation of dengue infection ^[6]. A primary antibody response is observed in individuals who are not immune to dengue and a secondary immune response is observed in patients who have had a previous dengue infection. In primary infection, IgM antibodies develop by 3-10 days after the onset of infection and reach its peak level ~2 weeks later. IgM generally decline to undetectable levels over the next 2-3 months ^[7, 8]. After the end of first week of infection, IgG is detectable at low titre and slowly increases, and persists for life. By contrast, during a secondary infection, high level of IgG antibodies is detectable even in the acute phase and rapidly rises over the following 2 weeks [9]. IgM antibodies may be detectable after 20 days of infection and are usually lower than in primary infections. IgA typically appears a day after IgM at a low level and decreases to undetectable level within 45 days following detection ^[10, 11, 12].

MP Diagnostics MULTISURE Dengue Ab/Ag Rapid Test is a reverse-flow immunochromatographic test and uses immobilized goat anti-human IgA, mouse anti-human IgM and IgG antibodies, and anti-dengue NS1 monoclonal antibodies for capturing IgA, IgM, IgG antibodies and NS1 dengue antigen in the human serum, plasma, finger pricked whole blood or whole blood with anti-coagulants. The presence of IgA, IgM, IgG antibodies against dengue virus and NS1 dengue antigen are detected by the colloidal gold-labeled dengue recombinant proteins and anti-dengue NS1 monoclonal antibody.

CHEMICAL AND BIOLOGICAL PRINCIPLES OF THE PROCEDURE

The **MP Diagnostic MULTISURE Dengue Ab/Ag Rapid Test kit** is a direct solid-phase immunochromatographic assay, based on MP Biomedicals' proprietary Reverse Flow technology (US Patent No.: 6,316,205) for simultaneous and differential detection of dengue antibodies and antigen in human serum, plasma, finger pricked whole blood or whole blood with anti-coagulants.

Goat anti-human IgA, mouse anti-human IgM and IgG, and dengue NS1 monoclonal antibodies are striped on the membrane as 4 separate test lines.

The dengue antibodies and/ or antigen in the test sample (serum, plasma, finger pricked whole blood or whole blood with anti-coagulants) form antibody-antigen complexes with anti-human antibodies and dengue NS1 monoclonal antibodies immobilized on nitrocellulose membrane in the test viewing window as the test sample migrates upward from the sample well.

The bound antibody-antigen complexes are subsequently detected by dengue antigens and dengue NS1 monoclonal antibody conjugates carried by chase buffer that flows downward giving a pink-purplish color. In addition, immobilized Biotinylated-BSA which can be recognized by colloidal gold-labeled goat anti-biotin is used as a control for proper function of the reagents.

DESCRIPTION OF SYMBOLS USED

The following are graphical symbols used in or found on MP Diagnostics' products and packaging. These symbols are the most common ones appearing on medical devices and their packaging.



KIT COMPONENTS

DEVICE	MP Diagnostics MULTISURE Dengue Ab/Ag Rapid Test Device	20 devices	
	Incorporated with anti-human and dengue NS1 monoclonal antibodies.		
	Packed in individually sealed pouch with a desiccant.		
	Store at 2°C - 28°C		
APPLICATOR	SAMPLE APPLICATOR	20	
	With marking of 20μ l and 25μ l.	pieces	
	20µ1 25µ1		
BUFFER	CHASE BUFFER	1 bottle	
	Contains sodium azide as preservative.	(5ml)	
	Store at 2°C - 28°C		
	INSTRUCTIONS FOR USE	1 copy	

HEALTH AND SAFETY INFORMATION

- 1. In case of an accident or contact with eyes, rinse immediately with plenty of water and seek medical advice.
- Consult a physician immediately in the event that contaminated materials are ingested or come in contact with open lacerations, or other breaks in the skin.
- 3. Wipe any spills of sera, plasma or blood promptly with 1% sodium hypochlorite solution.
- Autoclave all used and contaminated materials at 121°C, 15 p.s.i. for 30 minutes before disposal. Alternatively, decontaminate materials in 5% sodium hypochlorite solution for 30-60 minutes before disposal in biohazard waste-bags.

Pursuant to EC regulation 1272/2008 (CLP), hazardous components as classified and labelled as follows:

Component:	Chase Buffer
Signal Word:	Warning
Pictogram:	
Hazard Statements:	H319 Causes serious eye irritation.
Precautionary Statements:	P264 Wash hands thoroughly after handling. P280 Wear protective gloves/protective clothing/eye protection/ face protection.
Supplemental Statements:	EUH 210 Safety Data Sheet is available on request.
Contains:	1% Triton X-100

ANALYTICAL PRECAUTIONS

- 1. The sample applicator is for single use only. DO NOT re-use the sample applicator.
- Each sealed device is for single use only. DO NOT re-use the test device.
- 3. For Research use only.
- For Professional use only.
- 5. Wear protective clothing such as laboratory coats, disposable gloves and eye protection when samples are assayed.
- Handle all samples as if they contain infectious agents. Observe established precautions against microbiological hazards throughout all procedures and follow the standard procedures for proper disposal of samples.
- 7. Optimal assay performance requires **STRICT ADHERENCE** to the assay procedure described in this Instruction For Use. Deviations from the procedure may lead to aberrant results.
- 8. Do not interchange reagents between kit lots.
- 9. Do not use kit components beyond the expiry date printed on the kit box.
- The chase buffer reagent contains sodium azide as a preservative. Handle reagent carefully to avoid spills on work surface. Clean up spills with absorbent paper and water.
- 11. Reagents supplied should not present a hazard to health if used in accordance to the instructions stated.
- For best results, allow all reagents and samples to reach room temperature (25°C ± 3°C) before use.
- 13. Humidity and temperature can adversely affect results.
- 14. DO NOT use the test device if the seal of the pouch is broken.
- DO NOT use the test device if the pull tab is not fully inserted, i.e. prepulled.

STORAGE

- 1. Store the kit and its components at 2°C 28°C.
- 2. Do not freeze the kit and its components.
- 3. The test device must remain in the sealed pouch until use.
- 4. Do not use the device beyond the expiry date printed on the pouch.

MATERIAL REQUIRED BUT NOT PROVIDED

- 1. Timer
- Lancet
- 3. Alcohol swabs

SAMPLE COLLECTION

Whole blood can be collected in anti-coagulant containing tubes and used as outlined in the assay procedure immediately or stored at $2^{\circ}C$ to $8^{\circ}C$ for not more than 72 hours before use.

The presence of 6 anti-coagulants, Acid Citrate Dextrose (ACD), Citratephosphate-dextrose (CPD), Ethylenediaminetetraacetic acid (EDTA), Potassium Oxalate (K-Oxalate), Lithium Heparin (Li Hep) and Sodium Citrate (Na Citrate) were found to have no effects on the performance of MULTISURE Dengue Ab/Ag Rapid Test.

Serum / plasma samples should be stored at 2°C to 8°C if the test is to be run within 7 days of collection or frozen at -20°C or colder if the test is to be delayed for more than 7 days. Clear, non-hemolyzed samples are preferred. Lipemic, icteric or contaminated (particulate or bacterial) samples should be filtered (0.45 μ m) or centrifuged before testing.

Samples can be inactivated but this is not a requirement for optimal test performance.

Inactivate as follows:

- 1. Loosen cap of sample container.
- 2. Heat-inactivate sample at 56°C for 30 minutes in a water bath.
- 3. Allow sample to cool down before retightening cap.
- 4. Sample can be stored frozen until analysis.

Repeated freeze-thawing of the sample is not recommended. Do not use specimens (serum & plasma) with more than five (5) freeze-thaw cycles.

ASSAY PROCEDURE

IMPORTANT: Strict adherence to the assay procedure will ensure optimal performance. Deviations from the procedure may lead to aberrant results.

Note:

- Allow the kit to warm to room temperature $(25^{\circ}C \pm 3^{\circ}C)$ before running the assay.
- For best results, conduct the assay at room temperature ($25^{\circ}C \pm 3^{\circ}C$).
- 1. Open the pouch by tearing at the tear notch on the right side of the pouch to avoid accidental pulling of pull tab.
- Discard the device if pull tab is not fully inserted into test device. Refer to diagrams below.



- 3. Conduct the test immediately after removing the test device from the pouch.
- 4. Label the test device with the sample name.
- 5. Proceed with the assay procedures as shown in the diagrams below.

Assay Procedures for MULTISURE Dengue Ab/Ag Rapid Test



Do not read results after 25min.

QUALITY CONTROL

The control line is an internal procedure control. Absence of control line indicates incorrect assay technique used or faulty device. The procedure should be reviewed before repeating with a new device.

Example of test device

with all visible

test line

Positive and negative controls are not provided with this kit. However, it is recommended that controls are used as good laboratory practice to verify proper assay technique and performance.

INTERPRETATION OF RESULTS

1. Positive for Dengue

A test is dengue positive if Control Line (C) appears with any visible test lines (IgG, IgM, IgA, NS1).

- Presence of IgG indicates secondary or past infection.
- Presence of IgM and/or IgA indicates current infection.
- Presence of NS1 indicates acute dengue infection.

2. Negative for Dengue

A test is negative if Control line (C) appears and no visible test line(s). Retest in 3-5 days if dengue infection is suspected.

3. Invalid

A test is invalid if Control Line (C) is absent. The assay should be repeated using a new device.

LIMITATIONS OF THE PROCEDURE

The **MP Diagnostics MULTISURE Dengue Ab/Ag Rapid Test** is a qualitative *in vitro* immunochromatographic test to detect and differentiate IgA/ IgM/ IgG antibodies against dengue virus and NS1 dengue antigen in serum, plasma, finger pricked whole blood or whole blood with anti-coagulants. This test is not for determining the quantitative value in dengue antibodies/ antigen and in the diagnosis or prognosis of disease.

Optimal assay performance requires the strict adherence to the assay procedure described. Deviation from the procedure may lead to aberrant results.

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LIMITED EXPRESSED WARRANTY DISCLAIMER

The manufacturer makes no express warranty other than that the test kit will function as a Research Use Only assay within the specifications and limitations described in the product Instruction Manual when used in accordance with the instructions contained therein. The manufacturer disclaims any warranty expressed or implied, including such expressed or implied warranty with respect to merchantability, fitness for use or implied utility for any other purposes. The manufacturer is limited to either replacement of the product or refund of the purchase price of the product. The manufacturer shall not be liable to the purchaser or third parties for any damage, injury or economic loss howsoever caused by the product in the use or in the application thereof.

TECHNICAL PROBLEMS / COMPLAINTS

Should there be a technical problem / complaint, please do the following:

- 1. Note the kit lot number and the expiration date.
- 2. Retain the kits and the results that were obtained.
- 3. Contact the nearest MP Biomedicals office or your local distributor.



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