Malaria pan-LDH (PAN) Antigen Detection Test Device (Whole Blood) Instructions for Use
Catalog Number: B703C

A rapid test for the qualitative detection of Human Malaria antigen in whole blood.
For professional in vitro test use only.

Intended Use

For the rapid qualitative determination of one or more of the known Malaria species; Plasmodium falciparum, Plasmodium vivax, Plasmodium ovale, and/or Plasmodium malariae by detecting lactate dehydrogenase (LDH) in human blood. This test is an aid in the diagnosis of Malaria infection.

Summary

Malaria is a serious parasitic disease characterized by fever, chills, and anemia and is caused by a parasite that is transmitted from one human to another by the bite of infected Anopheles mosquitoes. There are four kinds of malaria that can infect humans: Plasmodium falciparum, Plasmodium vivax, Plasmodium ovale, and Plasmodium malariae. In humans, the parasites (called sporozoites) migrate to the liver where they mature and release another form, the merozoites. The disease now occurs in more than 90 countries worldwide, and it is estimated that there are over 500 million clinical cases and 2.7 million malaria-caused deaths per year. At the present, malaria is diagnosed by looking for the parasites in a drop of blood. Blood will be put onto a microscope slide and stained so that the parasites will be visible under a microscope.

The TELL ME FAST Malaria pan-LDH Antigen Detection Test Device (Whole Blood) contains a membrane strip, which is pre-coated with monoclonal antibodies on the test line region of the strip. When a Whole Blood specimen is applied at one end of the membrane and following the application of the assay buffer, it reacts with the antibody coated particles that have already been applied to the specimen pad. The mixture then migrates chromatographically towards the other end of the membrane and reacts with the monoclonal antibodies previously placed on the test line region. If the Whole Blood contains one or more of the four Malaria species, a colored line will appear in the test line region, showing a positive result. The absence of the colored line in the test region indicates a negative result therefore the whole blood does not contain detectable levels of any of the Malaria species. To serve as a procedural control, a colored line will always appear at the control line region if the test has been performed properly. This control line serves to validate the performance of the test.

Precautions

- For professional in vitro diagnostic use only. Do not use after expiration date.
- Do not eat, drink or smoke in the area where the specimens or kits are handled.
- Handle all specimens as if they contain infectious agents. Observe established precautions against microbiological hazards throughout testing and follow the standard procedures for proper disposal of specimens.
- Wear protective clothing such as laboratory coats, disposable gloves and eye protection when specimens are being tested.
- Humidity and temperature can adversely affect results.

Storage & Stability

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

Materials

Materials Provided:
- Test Device
- Assay Buffer
- Instructions for Use

Materials Not Provided:
- Calibrated pipette
- Lancet
- Timer
- Alcohol swab

Specimen Collection & Preparation

[Collection by venipuncture]
1) Collect whole blood into a collection tube (containing EDTA, citrate or heparin) by venipuncture.
2) If specimens are not immediately tested, they should be refrigerated at 2 ~ 8°C. For storage periods greater than three days, freezing is recommended. They should be brought to room temperature prior to use. Using the specimen after long-term storage of more than three days can cause non-specific reaction.
3) When stored at 2 ~ 8°C, the whole blood sample should be used within three days.

[Collection using a lancet]
1) Clean the area to be lanced with an alcohol swab.
2) Squeeze the end of the fingertip and pierce with a sterile lancet provided.
3) Wipe away the first drop of blood with sterile gauze or cotton.
4) Using a 5 μL calibrated dropper tube, while gently squeezing the tube, immerse the open end in the blood drop and then gently release the pressure to draw blood into the dropper.

Directions for Use:

Allow test device, buffer, specimen, and/or controls to equilibrate to room temperature (15-30°C) prior to testing.

1) Add 5 μl of whole blood into sample well [S], the small well.
2) Add two drops (80 μLs) of assay buffer

Manufactured by: Biocan Diagnostics Inc 160 Suite 309 North Vancouver BC Canada
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Interpretation of Results:

(Please refer to the illustrations)

1) The presence of two color bands indicates a positive result for Malaria. The pan-LDH present in the sample reacts with the anti-pan-LDH conjugate and moves through the test strip where the newly formed complex is captured by pan specific anti-pLDH.

2) Negative reaction
The presence of only one band in the control region of the result window indicates a negative result.

3) Invalid
The test is invalid if the control line does not appear. If this occurs, the test should be repeated using a new strip.

Limitations

1) The test procedure, precautions and interpretation of results for this test must be followed when testing.
2) Anti-coagulants such as heparin, EDTA, and citrate do not affect the test result.
3) This test kit detects lactate dehydrogenase in patient whole blood and is useful as a screening procedure of malaria diagnosis.
4) Do not mix reagent of different lots.
5) The test is limited to the detection of antigen to Malaria Plasmodium sp. Although the test is very accurate in detecting pan-LDH, a low incidence of false results can occur. Other clinically available tests are required if questionable results are obtained. As with all diagnostic tests, a definitive clinical diagnosis should not be based on the results of a single test, but should only be made by the physician after all clinical and laboratory findings have been evaluated.

Bibliography


