



AccuBiotech Co., Ltd.

ACCU-TELL®

**Malaria p.f./p.v./pan Cassette
(Whole Blood)**

For in vitro diagnostic Use only

For Whole Blood Sample

This package insert is applied to the below products:

Catalog No. Product Name

ABT-IDT-B236 Malaria p.f./p.v./pan Cassette (Whole Blood)

A rapid test for the qualitative detection of circulating antigens of *P. falciparum* (P.f.), *P. vivax* (P.v.), *P. ovale* (P.o.), and *P. malariae* (P.m.) in whole blood

For professional in vitro diagnostic use only

INTENDED USE

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) is a rapid chromatographic immunoassay for the qualitative detection of four kinds of circulating plasmodium falciparum (*P. falciparum* (P.f.), *P. vivax* (P.v.), *P. ovale* (P.o.), and *P. malariae* (P.m.) in whole blood.

SUMMARY

Malaria is caused by a protozoan which invades human red blood cells.¹ Malaria is one of the world's most prevalent diseases. According to the WHO, the worldwide prevalence of the disease is estimated to be 300-500 million cases and over 1 million deaths each year. Most of these victims are infants, young children. Over half of the world's population lives in malarious areas. Microscopic analysis of appropriately stained thick and thin blood smears has been the standard diagnostic technique for identifying malaria infections for more than a century.² The technique is capable of accurate and reliable diagnosis when performed by skilled microscopists using defined protocols. The skill of the microscopist and use of proven and defined procedures, frequently present the greatest obstacles to fully achieving the potential accuracy of microscopic diagnosis. Although there is a logistical burden associated with performing a time-intensive, labor-intensive, and equipment-intensive procedure such as diagnostic microscopy, it is the training required to establish and sustain competent performance of microscopy that poses the greatest difficulty in employing this diagnostic technology.

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) is a rapid test to qualitatively detect the presence of *P. falciparum* - specific HRP-II and four kinds of circulating plasmodium falciparum (*P. falciparum* (P.f.), *P. vivax* (P.v.), *P. ovale* (P.o.), and *P. malariae* (P.m.)). The test utilizes colloid gold conjugate to selectively detect P.f-specific and Pan-malarial antigens (P.f., P.v., P.o. and P.m.) in whole blood.

PRINCIPLE

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) is a qualitative, membrane based immunoassay for the detection of P.f., P.v., P.o. and P.m. antigens in whole blood. The membrane is pre-coated with anti-HRP-II antibodies, anti-P.vivax specific LDH antibodies and anti-Pan specific LDH antibodies. During testing, the whole blood specimen reacts with the dye conjugate, which has been pre-coated on the test cassette. The mixture then migrates upward on the membrane by capillary action, reacts with anti-Histidine-Rich Protein II (HRP-II) antibodies on the membrane on P.f. Test Line region, with anti-P.vivax specific LDH antibodies on the membrane on P.v. Test Line region and with anti-Pan specific LDH antibodies on the membrane on Pan Line region. If the specimen contains HRP-II or P. vivax specific LDH or Pan specific LDH or all, a colored line will appear in P.f. line region or P.v. line region or Pan line region or three colored lines will appear in P.f. line region, P.v. line region and Pan line region.

The absence of the colored lines in P.f. line region or P.v. line region or Pan line region indicates that the specimen does not contain HRP-II and/or P.vivax specific LDH and/or Pan specific LDH. To serve as a procedure control, a colored line will always appear in the control line region indicating that proper volume of specimen has been added and membrane wicking has occurred.

REAGENTS

The test cassette contains anti-HRP-II of *Plasmodium falciparum* antibodies conjugated gold, anti- *P.vivax* specific LDH antibodies conjugated gold and anti-Pan specific LDH antibodies conjugated gold and anti-HRP-II antibodies, anti-P.vivax specific LDH antibodies and anti-Pan specific LDH antibodies coated on the membrane.

PRECAUTIONS

- For professional in vitro diagnostic use only. Do not use after expiration date.
- For whole blood specimen use only. Do not use other specimens.
- 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 all procedures 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 assayed.
- The used tests, specimens and potentially contaminated materials should be discarded according to the local regulations.
- Humidity and temperature can adversely affect results.
- Do not exchange or mix buffer and test cassettes from kits of different lot numbers.
- Caution must be taken at the time of specimen collection. Inadequate volume of specimen may lead to lower sensitivity.
- Be sure to add sufficient buffer to the cassette's sample well. Invalid result may occur if inadequate buffer is added.

STORAGE AND STABILITY

The kit can be stored at room temperature or refrigerated (2-30°C). The test cassette is stable through the expiration date printed on the sealed pouch. The uncovered buffer could be stored at room temperature or refrigerated (2-30°C) for 1.5 months at least. The test cassette must remain in the sealed pouch until use. **DO NOT FREEZE.** Do not use beyond the expiration date.

SPECIMEN COLLECTION AND PREPARATION

- ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) can be performed using whole blood.
- Both Fingerstick Whole Blood and Venipuncture Whole Blood can be used.
- **To collect Fingerstick Whole Blood specimens:**
- Wash the patient's hand with soap and warm water or clean with an alcohol swab. Allow to dry.
- Massage the hand without touching the puncture site by rubbing down the hand towards the fingertip of the middle or ring finger
- Puncture the skin with a sterile lancet. Wipe away the first sign of blood.
- Gently rub the hand from wrist to palm to finger to form a rounded drop of blood over the puncture site.
- Testing should be performed immediately after specimen collection. Do not leave the specimens at room temperature for prolonged periods. Whole blood collected by venipuncture should be stored at 2-8°C if the test is to be run within 2 days of collection. For long term storage, specimens should be kept below -20°C. Whole blood collected by fingerstick should be tested immediately.
- 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 for more than three times.

- If specimens are to be shipped, they should be packed in compliance with federal regulations covering the transportation of etiologic agents.

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Version #: HBT-Effective

5221404

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood)

Date: 2022-03 AccuBiotech Co., Ltd.

MATERIALS Materials Provided

Test Cassettes
Disposable specimen droppers
Buffer
Package insert

Materials Required But Not Provided

Pipette and disposable tips (optional)
Specimen collection containers
Lancets (for fingerstick whole blood only)
Timer

TEST PROCEDURE

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

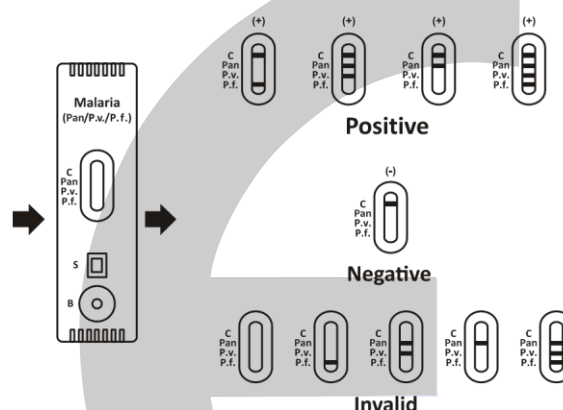
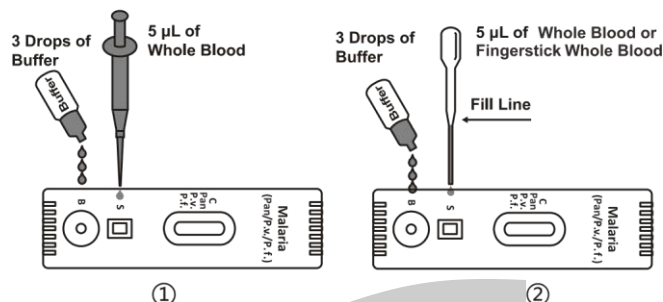
- Bring the pouch to room temperature before opening it. Remove the test cassette from the sealed pouch and use it within one hour.
- Place the cassette on a clean and level surface.

For Whole Blood specimen:

- Use a pipette: To transfer 5µL of whole blood to the specimen well, then add 3 drops of buffer (approximately 180µL).
- Use a disposal specimen dropper: Hold the dropper vertically, draw the specimen up to the Fill Line as shown in illustration below (approximately 5µL). Transfer the specimen to the specimen well, then add 3 drops of buffer (approximately 180µL), and start the timer.

For Fingerstick Whole Blood

- Use a disposal specimen dropper: Hold the dropper vertically, draw the specimen up to the Fill Line as shown in illustration below (approximately 5µL). Transfer the specimen to the specimen well, then add 3 drops of buffer (approximately 180µL), and start the timer.
- Wait for the colored line(s) to appear. Read results at **10 minutes**. Do not interpret the result after 20 minutes.



INTERPRETATION OF RESULTS

(Please refer to the illustration above)

POSITIVE: * Two or Three or Four distinct colored lines appear.

P.f. infection: One colored line should be in the control line region (C), and one colored line appears in P.f. line region.

P.v. infection: One colored line should be in the control line region (C), one colored line appears in P.v. line region, and/or one colored line appears in Pan line region. If the concentration of P.v. is too low, maybe no line appears in Pan line region.

P.o. and/or P.m. infection: One colored line should be in the control line region (C), and one colored line appears in Pan line region.

P.f. and P.o. and/or P.m. infection, P.f. and/or P.o. and P.m. infection: One colored line should be in the control line region (C), one colored line appears in P.f. line region, and one colored line appears in Pan line region.

P.v. and P.o. and/or P.m. infection, P.v. and/or P.o. and P.m. infection: One colored line should be in the control line region (C), one colored line appears in P.v. line region, and one colored line appears in Pan line region.

Mixed infection: One colored line should be in the control line region (C), one colored line appears in Pan line region, one colored line appears in P.v. line region and One colored line appears in P.f. region.

***NOTE:** The color intensity of Pan, P.v. and/or P.f. test lines may vary depending on the concentration of antigens, viz., HRP-II, P. vivax specific LDH or Pan specific LDH present in the specimen. **NEGATIVE:**



One colored line appears in the control line region (C). No line appears in the test line region (T).

INVALID: Control line fails to appear. Insufficient specimen volume is 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 cassette 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 procedural control. It confirms adequate membrane wicking.

Control standards are not supplied with this kit; however, 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.

LIMITATIONS

1. ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) is for in vitro diagnostic use only. This test should be used for the detection of P.f., P.v., P.o., P.m. antigens in whole blood specimens only. Neither the quantitative value nor the rate of increase in P.f., P.v., P.o., and P.m. concentration can be determined by this qualitative test.

2. ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) will only indicate the presence of antigens of Plasmodium sp. (P.f., P.v., P.o., P.m.) in the specimen and should not be used as the sole criterion for the diagnosis of malaria infection.
3. As with all diagnostic tests, all results must be interpreted together with other clinical information available to the physician.
4. 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 malaria infection.

EXPECTED VALUES

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) has been compared with traditional thick and thin blood films microscopic analysis. The correlation between the two systems is over 99.0%.

PERFORMANCE CHARACTERISTICS Sensitivity

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) has been tested with microscopy on clinical samples. The results show that the overall sensitivity of P. falciparum and P. vivax were >99.9% and 98.0% when compared to results obtained with microscopy. **Specificity**

ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) uses antibodies that are highly specific to Malaria P.f.-specific and Pan-

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malarial antigens in whole blood. The results show that the specificity malaria parasite in human blood samples. Am J Trop Med of ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) Hyp,1999, Feb: 60(2):173-2. is >99.9%, when compared to results obtained with microscopy.

Method	Results	Microscopy		Total Results
		Positive	Negative	
ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood)	Positive	50*	82**	132
	Negative	1	0	432
Total Results		51	82	432

GLOSSARY OF SYMBOLS

REF	Catalog number		Temperature limitation
	Consult instructions for use	LOT	Batch code
IVD	In vitro diagnostic medical device		Use by
	Manufacturer		Do not reuse

Comment : Blood Samples infected by Plasmodium falciparum (n = 82), Plasmodium vivax (n = 51) were included, as well as 432 malaria negative samples to be confirmed with microscopy.

Note: * There was one P. vivax sample to show a pan line and a P.f. line.

**There were two P. falciparum samples that they both showed a pan line and a P.f. line.

The comparison for Pan line has been only done with blood specimens positive with Plasmodium vivax specimen. The claims for Pan lines are based on scientific findings that Pan specific LDH is found in other malarial parasites including Plasmodium ovale and Plasmodium malariae.

Relative Sensitivity for P.f.-specific antigens:

82/82>99.9 % (95%CI***: 96.4%~100.0%)

Relative Sensitivity for Pan-malarial antigens:

50/51= 98.0% (95%CI***: 89.6%~100.0%) **Relative**

Specificity:

432/432>99.9% (95%CI***: 99.3%-100.0%)*

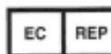
Accuracy: (50+82+432)/(82+51+432)=564/565=99.8%(95%CI***: 99.0%-100.0%)

***Confidence Intervals



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Minimum Detection Level

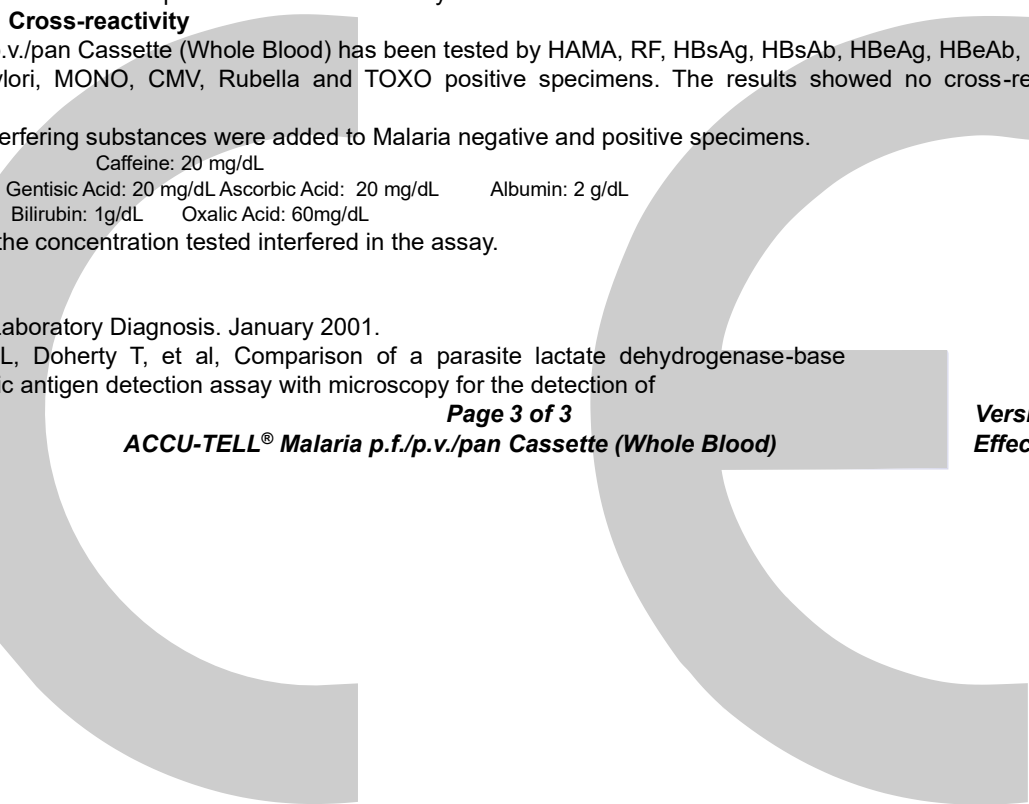
Type	Parasites/□L
P. falciparum	200
Plasmodium non-falciparum species (P. vivax)	1500

Precision Intra-Assay

Within-run precision has been determined by using 15 replicates of four specimens: a negative, a P.f. positive, a P.v. positive and an P.f./P.v. dual positive. The specimens were correctly identified >99% of the time.

Inter-Assay

Between-run precision has been determined by 15 independent assays on the same four specimens: negative, a P.f. positive, a P.v. positive and an P.f./P.v. dual positive. Three different lots of ACCUTELL® Malaria p.f./p.v./pan Cassette (Whole Blood) have been tested using these specimens. The specimens were correctly identified >99% of the time.



Cross-reactivity
ACCU-TELL® Malaria p.f./p.v./pan Cassette (Whole Blood) has been tested by HAMA, RF, HBsAg, HBsAb, HBeAg, HBeAb, HBcAb, Syphilis, HIV, HCV, H. Pylori, MONO, CMV, Rubella and TOXO positive specimens. The results showed no cross-reactivity.

Interfering Substances

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

Acetaminophen: 20 mg/dL Caffeine: 20 mg/dL
Acetylsalicylic Acid: 20 mg/dL Gentisic Acid: 20 mg/dL Ascorbic Acid: 20 mg/dL Albumin: 2 g/dL
Creatin: 200 mg/dL Bilirubin: 1g/dL Oxalic Acid: 60mg/dL

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

BIBLIOGRAPHY

1. Bill MaConell, Malaria Laboratory Diagnosis. January 2001.
2. Cooke AH, Chiodini PL, Doherty T, et al, Comparison of a parasite lactate dehydrogenase-base immunochromatographic antigen detection assay with microscopy for the detection of