WHO Prequalification of In Vitro Diagnostics PUBLIC REPORT

Product: First Response HIV1+2/Syphilis Combo Card Test WHO reference number: PQDx 0364-010-00

First Response HIV1+2/Syphilis Combo Card Test with product codes **I20FRC25**, **I20FRC30**, **I20FRC50**, **I20FRC60**, **I20FRC100**, **and I20FRC25-SA**, manufactured by **Premier Medical Corporation Private Limited**, **Rest of World regulatory version**, was accepted for the WHO list of prequalified in vitro diagnostics and was listed on 24 June 2019.

Summary of WHO Prequalification Assessment for First Response HIV1+2/Syphilis Combo Card Test

	Date	Outcome
Prequalification listing	24 June 2019	listed
Dossier assessment	2 May 2019	MR
Site inspection(s) of the	15-17 October 2022	MR
quality management system		
Product performance	Q1 of 2018	MR
evaluation		

MR: Meets Requirements

This public report has since been amended. Amendments may have arisen because of changes to the prequalified product for which WHO has been notified and has undertaken a review. Amendments to the report are summarized in the following table, and details of each amendment are provided below.

Version	Summary of amendment	Date of report amendment
2.0	A new specimen transfer device with a 20 μl marking line was introduced to make it more user-friendly.	6 February 2020
3.0	Introduction of new pack size with auto safety lancet, with product code I20FRC25-SA.	9 August 2023
4.0	Addition of performance evaluation results for HIV subtypes.	27 March 2024

Intended use

According to the claim of intended use from Premier Medical Corporation Private Limited, "First Response HIV 1+2 / Syphilis Combo Card Test is intended for use by healthcare professionals and trained users. It is a rapid, qualitative screening, in vitro diagnostic test for the detection of antibodies (IgG & IgM) specific to HIV (type 1 & 2) and Treponema pallidum in human serum, plasma or venous and capillary whole blood. The test can be used as an aid in diagnosis of HIV and/or Syphilis. The product is intended to be used for symptomatic, asymptomatic as well as pregnant women population. The test kit is not automated and does not require any additional instrument. Reactive specimens should be confirmed further with ELISA, Western Blot or TPHA ".

Assay description

According to the claim of assay description from Premier Medical Corporation Private Limited "First Response HIV 1+2 / Syphilis Combo Card Test is based on the principle of immunochromatography for qualitative detection of antibodies (IgG & IgM) specific for HIV 1&2 and/or Syphilis. The nitrocellulose membrane is coated with cocktail of recombinant antigen for HIV 1 (gp41) and HIV 2 (gp36) at test line "HIV" and Recombinant TP antigen (P47, P45, P17, P15) specific for Treponema pallidum at the test line "Syp" and control reagent coated at control line "C". When a serum or plasma or whole blood specimen is applied to the specimen well of test device, the cocktail of recombinant HIV 1+2 (gp41 & gp36) antigen - colloidal gold conjugate (CGC) & recombinant Treponema pallidum antigens colloidal gold conjugate, will react with HIV and/or Syphilis specific antibodies, if present in the specimen. The antibody-CGC antigen complex and assay buffer move along the membrane chromatographically to the test regions and form a visible line as the antigen-antibody-CGC antigen complex forms with high degree of sensitivity and specificity.

If the specimen contains antibodies to Treponema pallidum, the colored line will appear in the test area at test line "Syp", corresponding to Syphilis line. If the specimen contains antibodies to HIV 1 and/or 2, the colored line will appear in the test area at test line "HIV", corresponding to HIV 1+2 line"

Component	25 tests (product code I20FRC25)	30 tests (product code I20FRC30)	50 tests (product code I20FRC50)	60 tests (product code I20FRC60)	100 tests (product code I20FRC100)	25 tests (product code I20FRC25-SA)
Test device pouch containing: 1 test device, 1 desiccant	25	30	50	60	100	25
Specimen transfer device	25	30	50	60	100	25

Test kit contents

Assay buffer	1 of 2.5 ml	1 of 2.5 ml	2 of 2.5	4 of 2.5	4 of 2.5 ml	1 of 2.5 ml
bottle			ml	ml		
Sterile twist	25	30	50	60	100	\
lancets						
Auto Safety	\	\	\	\	\	25
Lancet						
Alcohol swabs	25	30	50	60	100	25
Instructions for	1	1	1	1	2	1
use						

Items required but not provided

- New pair of disposable gloves and face mask.
- Permanent Marker pen and timer
- Extra lancets and alcohol swabs, if needed
- Sharp disposable box and biohazardous waste container
- Venipuncture blood collection kit (if whole blood is collected by venipuncture
- Sterile gauze pads

Storage

The test kit should be stored at 4-30 °C.

Shelf-life upon manufacture

24 months.

Warnings/limitations

Refer to the current version of the manufacturer's instructions for use.

Prioritization for prequalification

Based on the established eligibility criteria, the First Response HIV 1+2 / Syphilis Combo Card Test was given priority for WHO prequalification assessment.

Dossier assessment

Premier Medical Corporation Private Limited submitted a product dossier for the First Response HIV1+2/Syphilis Combo Card Test as per the "Instructions for compilation of a product dossier" (PQDx_018 version 3). The information (data and documentation)

submitted in the product dossier was reviewed by WHO staff and external technical experts (assessors) appointed by WHO.

The manufacturer's responses to the nonconformities found during dossier screening and assessment findings were accepted on 02 May 2019.

Based on the product dossier screening and assessment findings, the product dossier for the First Response HIV1+2/Syphilis Combo Card Test meets WHO prequalification requirements.

Manufacturing site inspection

An onsite inspection of Premier Medical Corporation Ltd., at A1-302 and 3704-05, GIDC, Sarigam INA, 396155 Gujarat, India, was conducted from the 15th to the 17th of October 2022. At the time of considering the product application for Prequalification, the Manufacturer of the product had a well-established quality management system and manufacturing practices in place that would support the manufacture of a product of consistent quality. Routine inspections of the Manufacturing site will be conducted with copies of the WHO Public Inspection Report (WHOPIR) published on the WHO Prequalification web page as per Resolution WHA57.14 of the World Health Assembly. Note that a WHOPIR reflects the information on the most current assessment performed at a manufacturing site for in vitro diagnostic products and summarises the assessment findings.

https://extranet.who.int/pqweb/vitro-diagnostics/who-public-inspection-reports

All published WHOPIRs are with the agreement of the manufacturer. The onsite inspection was accepted on 25 September 2023.

Product performance evaluation

First Response HIV 1+ 2/Syphilis Combo card test (Premier Medical Corporation Private Limited) is a single-use, rapid, qualitative lateral flow immunochromatography assay for the detection of HIV-1/2 and Syphilis antibodies in human serum/plasma, whole blood (finger stick, EDTA, heparin or sodium citrate). A volume of 20 μ L of specimen is needed to perform the assay. This type of assay requires nosophisticated equipment and can, therefore, be performed in laboratories with limited facilities and non-laboratory settings. Reading of the results can be done visually, i.e. subjectively read.

First Response HIV 1+ 2/Syphilis Combo card test (Premier Medical Corporation Private Limited) was evaluated by WHO in the first quarter of 2018 at the Institute of Tropical Medicine, Belgium, using serum/plasma specimens.

In this limited evaluation on a panel of 400 clinically-derived specimens, compared to the reference assays (HIV reference algorithm: Vironostika HIV Ag/Ab [bioMérieux] EIA and Enzygnost Anti-HIV 1/2 Plus [Siemens Healthcare Diagnostics] EIA or Genscreen HIV-1/2 Version 2 [Bio-Rad]; followed by INNO-LIA HIV I/II Score [Fujirebio Inc.]; Syphilis reference algorithm: Vitros Syphilis TPA Assay [Ortho Clinical Diagnostics], followed by SERODIA-TP.PA [Fujirebio Inc.]), the following results were obtained:

Performance characteristics in comparison with an agreed reference standard								
	HIV-:	1/2	Syphilis					
	Initial (95% CI) Final (95% CI)		Initial (95% CI)	Final (95% CI)				
Sensitivity %	100%	100%	99.0%	99.0%				
(N=200)	(98.2% - 100%)	(98.2% - 100%)	(96.4% - 99.9)	(96.4% - 99.9%)				
Specificity %	99.0%	99.0% 99.5%		100%				
(N=200)	(96.4% - 99.9%)	(97.2% - 100%)	(96.4% - 99.9%)	(98.2% - 100%)				
Invalid rate	0%							
Inter-reader variability	0.5	%	0.3%					

In addition, analytical performance characteristics were assessed using commercially available panels and the following results were obtained:

Additional performance cha	Additional performance characteristics								
	HIV-1/2	Syphilis							
Sensitivity during	Seroconversion sensitivity	Seroconversion sensitivity index							
seroconversion in	index of 0, therefore detection	of -1, therefore detection is 1							
comparison with a	is 0 specimens earlier/later	specimen earlier than the							
benchmark assay	than the benchmark assay	benchmark assay (Vitros Syphilis							
	(Enzygnost Anti-HIV 1/2 Plus)	TPA Assay) on one							
	on average on 8 seroconversion	seroconversion panel							
	panels								
Analytical sensitivity on a	25 of 25 specimens were	17 of 17 specimens were							
mixed titer panel in	correctly classified	correctly classified							
comparison with an									
agreed reference standard									
Analytical sensitivity on	All 6 HIV subtypes/groups in	The 1 st International Standard for							
WHO reference	the 1 st International Reference	human syphilitic plasma IgG							
preparation panels	Panel for anti-HIV (NIBSC code	(NIBSC code 05/122) was							
	02/210) were detected	detected							
Lot to lot variation on a	Acceptable	Acceptable – except for one 2-							
dilution panel		dilution difference in one of 10							
		dilution panels							

Key operational characteristics	
Validated specimen types	Serum, plasma (EDTA, heparin or sodium citrate), venous whole blood, capillary whole blood
Number of steps	3 without precision required
Time to result	15 minutes
Endpoint stability	10 minutes (do not interpret after 25 minutes after addition of buffer)
Internal QC	Yes, control line on the test device
In-use stability of reagents	The assay buffer (opened & unopened) & the unopened test device are stable until the expiry date printed on the label when stored at 4-30°C.

Labelling

- 1. Labels
- 2. Instructions for use

- 1. Labels
- 1.1 Sterile safety lancet label (part of product codes I20FRC25, I20FRC30, I20FRC50, I20FRC60, and I20FRC100)



1.2 Sterile Pressure Activated Safety Lancet (part of product code I20FRC25-SA)



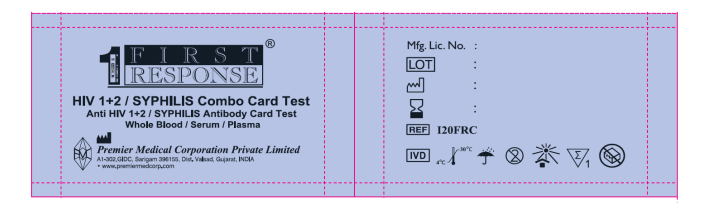
1.3 Alcohol swab labels

<text></text>	<text><text><text><text></text></text></text></text>
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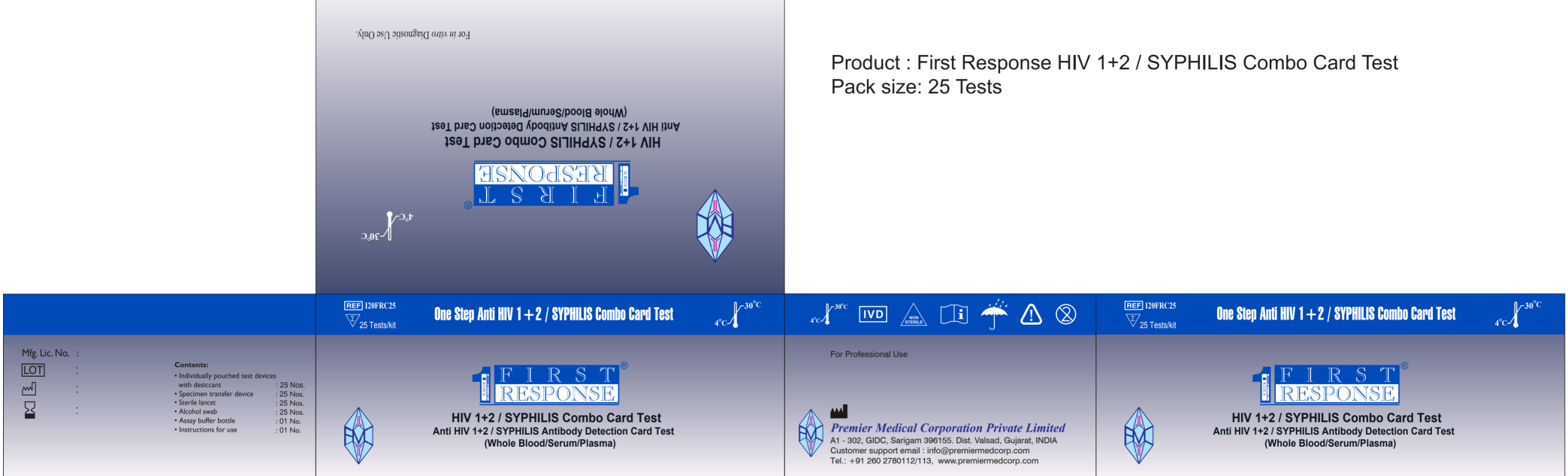
1.4 Assay buffer label

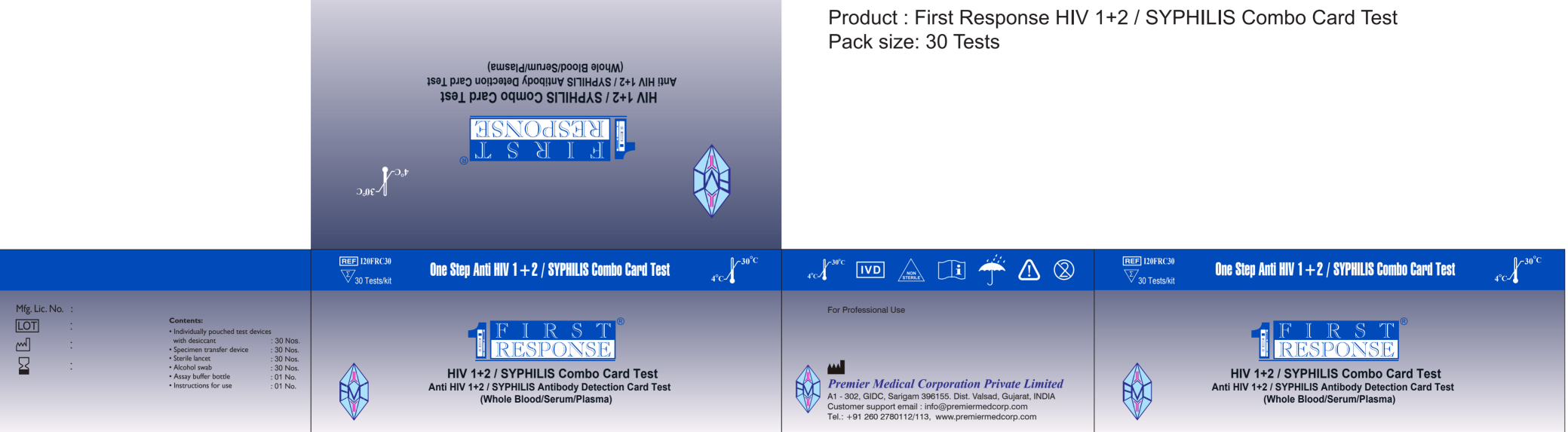


1.5 Aluminium pouch labels

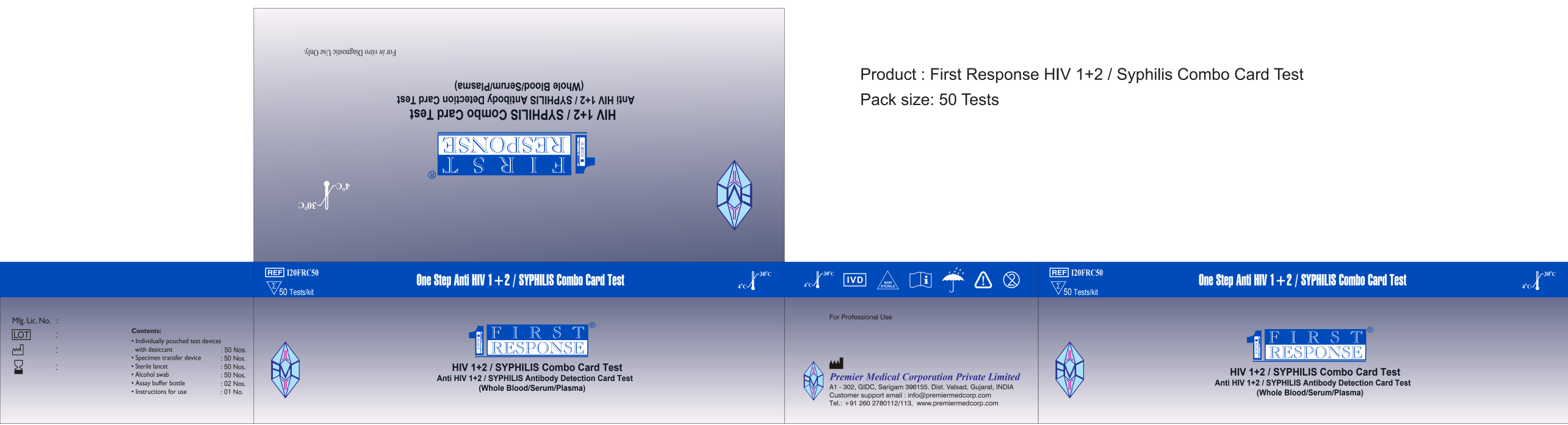


1.6 Outside box artwork and labels





For in vitro Diagnostic Use Only.







HIV 1+2 / SYPHILIS Combo Card Test Anti HIV 1+2 / SYPHILIS Antibody Detection Card Test (Whole Blood/Serum/Plasma)

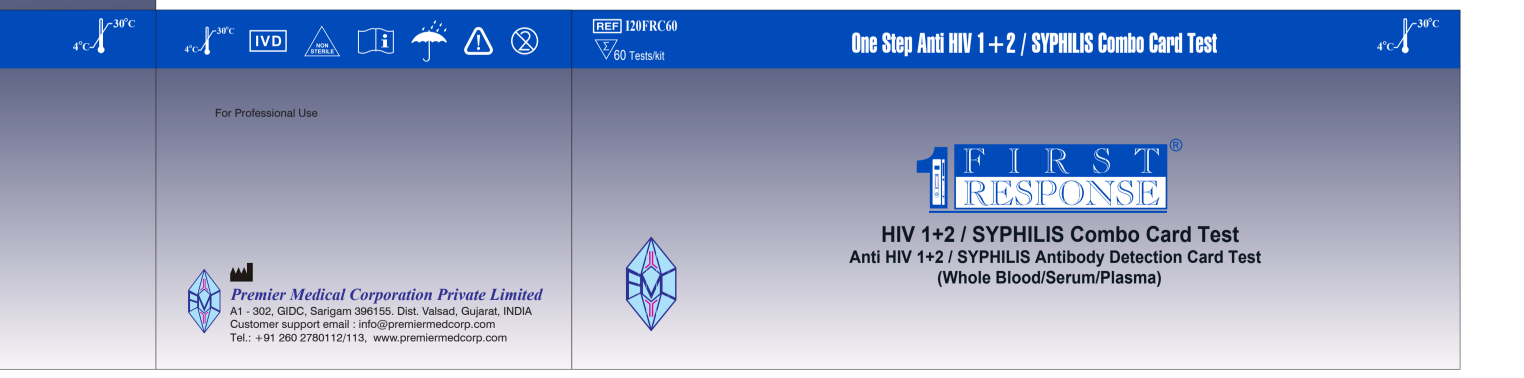


(Whole Blood/Serum/Plasma) Anti HIV 1+2 / SYPHILIS Antibody Detection Card Test HIV 1+2 / SYPHILIS Combo Card Test

For in vitro Diagnostic Use Only.

Product : First Response HIV 1+2 / SYPHILIS Combo Card Test Pack size: 60 Tests







Contents:

Individually pouched test devices

with desiccant

Sterile lancet

Alcohol swab

• Assay buffer bottle

Instructions for use

Specimen transfer device

- ∑ 100 Tests/kit

: 100 Nos.

: 100 Nos.

: 100 Nos.

: 100 Nos.

: 04 Nos.

: 02 Nos.





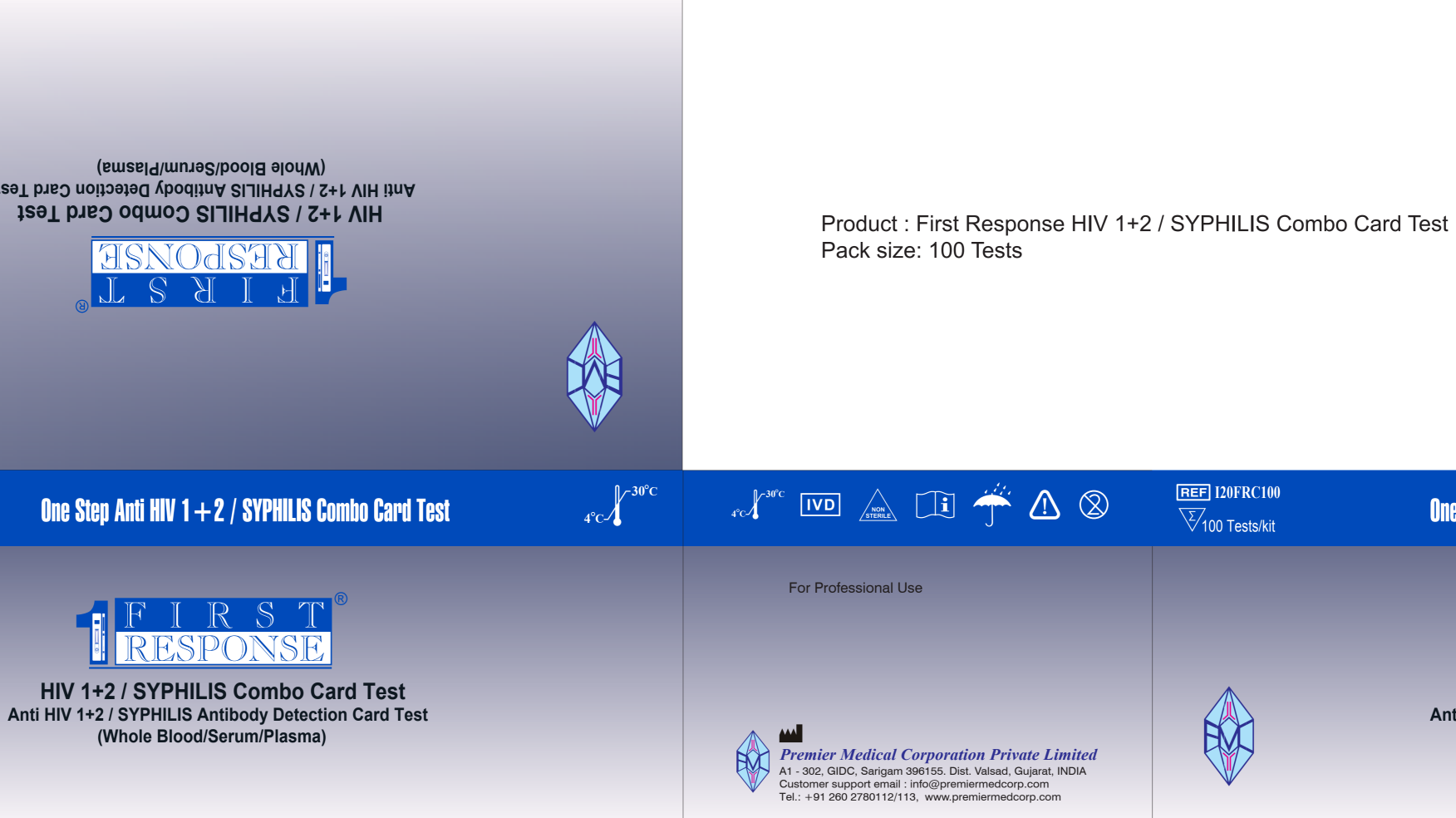
REF I20FRC100



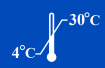
(Whole Blood/Serum/Plasma) Anti HIV 1+2 / SYPHILIS Antibody Detection Card Test HIV 1+2 / SYPHILIS Combo Card Test

For in vitro Diagnostic Use Only.



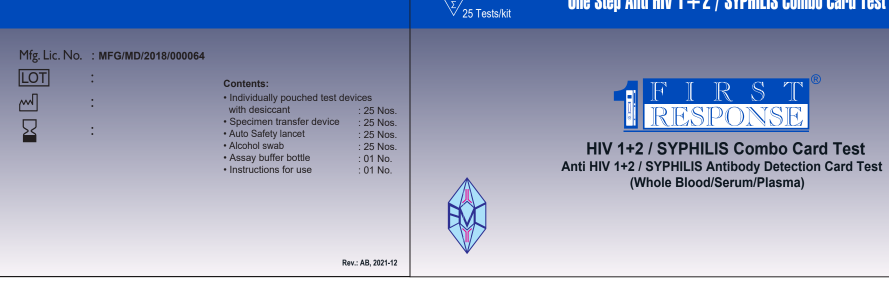


One Step Anti HIV 1 + 2 / SYPHILIS Combo Card Test





HIV 1+2 / SYPHILIS Combo Card Test Anti HIV 1+2 / SYPHILIS Antibody Detection Card Test (Whole Blood/Serum/Plasma)





One Step Anti HIV 1 + 2 / SYPHILIS Combo Card Test

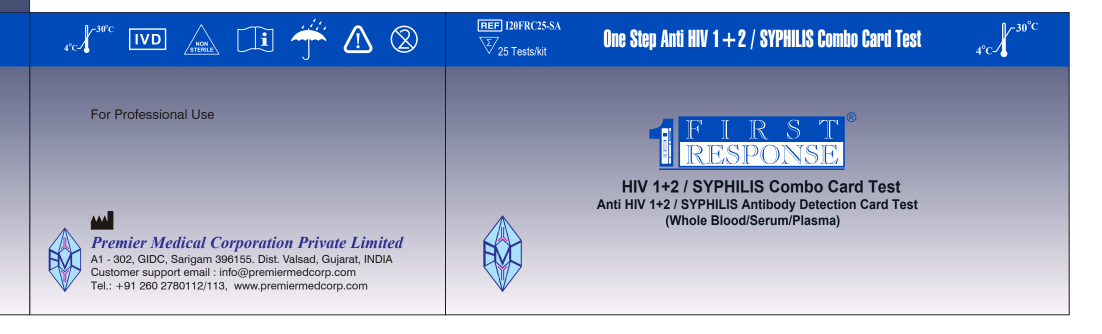






(Whole Blood/Serum/Plasma) Anti HIV 1+2 / SYPHILIS Antibody Detection Card Test HIV 1+2 / SYPHILIS Combo Card Test

For in vitro Diagnostic Use Only.



2. Instructions for use¹

¹ English version of the IFU was the one that was assessed by WHO. It is the responsibility of the manufacturer to ensure correct translation into other languages

- 6) Do not use the test device if the pouch is not intact.
- 7) Do not use the sterile twist lancet if the seal is broken (Refer specimen collection section)
- 8) Do not use the test device if the desiccant color has changed from orange to green.
- 9) Do not smoke, eat or drink while handling specimens and performing a test.
- 10) Do not re-use the test device, alcohol swab, sterile twist lancet, and specimen transfer device as these are intended for single use only.
- 11) Perform the test by using kit assay buffer, any other buffer or fluid will invalidate the test results
- 12) Do not allow the tip of assay buffer bottle to touch specimen well as it may contaminates the assay buffer.
- 13) Do not use the test device and assay buffer beyond the date of expiry.
- 14) Do not eat the desiccant.
- 15) Do not use any other specimen other than human Whole blood/Serum/Plasma. Do not mix and interchange different specimens.

Specimen Collection

- 1) Venous blood collection: Collect the Whole blood in the collection tubes containing anticoagulants like EDTA, Heparin, Sodium citrate or ACD by venipuncture
- 2) Plasma collection: Collect the Whole blood in the collection tubes containing anticoagulants like EDTA, Heparin, Sodium citrate or ACD by venipuncture and centrifuge it at 3000 g for 10-15 minutes to obtain Plasma.
- 3) Serum collection: Collect Whole blood in the collection tubes without having any anticoagulants by venipuncture. Keep it in standing position for 30 minutes and centrifuge it at 3000 g for 10-15 minutes to obtain serum.

4) Capillary whole blood specimen collection:

- Wear gloves and massage the fingertip gently. It will help to obtain a round drop of blood
- Wipe the complete fingertip with the alcohol swab provided and wait until the fingertip dried completely.



Verify the seal before detaching the cap. Sidelock confirms integrity of sterile twist lancet. Detach the protective cap of the sterile twist lancet. Squeeze the fingertip then prick the lateral side (avoid callus) of the fingertip with sterile twist lancet provided. Safely dispose of the used sterile twist lancet in sharps container immidiately after use...

- Wipe the first drop of the blood using sterile gauze. Without pressing too hard, gently squeeze fingertip once again to obtain the second drop of blood (~40-50 µl).
- Take the specimen transfer device provided and hold it vertically. Gently squeeze the bulb of specimen transfer device and immerse open end in the center of a blood drop and release the bulb slowly to draw up the blood up to the 20 µl marking line on the specimen transfer device.



-20µl marking • Do not use the specimen transfer device having no marking. After completion of specimen collection, take the sterile gauze and apply pressure to the wound site to stop the bleeding. The specimen transfer device is for single use only.

Note: Sterile twist lancet is for single use only. Do not share used sterile twist lancet with another person. Dispose of used sterile twist lancet in sharp box and alcohol swab in biohazard waste container immediately after use.

Do not use expired sterile twist lancet. Use of any expired sterile twist lancet may cause infections at the punctured skin due to the expiry of its sterility. Use new sterile twist lancet, alcohol swab and specimen transfer device and choose a different puncture site, if another finger pricking is required.

Specimen storage

1) Venous whole blood specimen should be used for testing immediately (within 1 hour) or shall be stored at 2-8°C for up to 72 hours (3 days). Do not use whole blood specimen stored for more than 3 days, it can cause a non-specific reaction. Do not freeze whole blood specimens.

Note: Mix the whole blood specimens in the tube by inverting the tube 3 or 4 times before use.

- 2) If serum or plasma specimens are not immediately tested, then they should be refrigerated at 2-8°C. For storage period greater than 72 hours (3 days), freezing at <-20°C is recommended up to 4 months.
- 3) Venous whole blood, serum and plasma specimens stored at 2-8 °C must be brought to room temperature before use. Serum or plasma specimens stored at <-20°C must be thawed at 15 to 25°C. Avoid more than 2 freeze-thaw cycles.

4) Serum or plasma specimens containing precipitate may yield inconsistent test results. Such specimens must be centrifuged at 5000 g for 10 minutes and then use clear supernatants for testing.

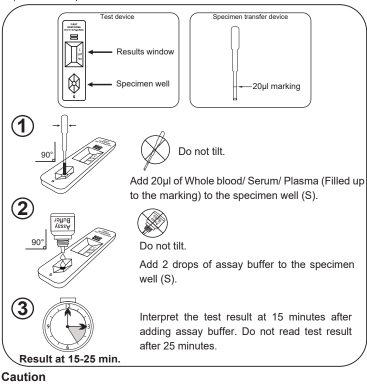
Test Procedure

- 1) Ensure that the test device & other components are at room temperature (15°C to 30°C) before starting the procedure.
- 2) Open the device pouch, take out the test device from aluminum pouch. Do not use the test device if the desiccant color has changed from orange to green.
- 3) Label the test device with the patient identification number. Place the test device on a flat, clean and dry surface.
- 4) Take out the specimen transfer device from plastic bag provided inside the kit. Gently squeeze the bulb of specimen transfer device and immerse the open end in the specimen and release the bulb slowly to draw up the serum/plasma/ capillary or venous whole blood up to 20µl marking line on the specimen transfer device.
- 5) Gently wipe away the excess specimen from the outer surface of the specimen transfer device with tissue paper before dispensing the specimen into the specimen well
- 6) Gently squeeze the bulb of specimen transfer device to add 20 µl of venous or capillary whole blood/ serum/ plasma to the specimen well by gently touching the tips of the specimen transfer device to the sample pad.

Caution: Dispose of used specimen transfer device and tissue paper as biohazard waste immediately after use

- 7) Hold the assay buffer bottle vertically and add two drops of assay buffer to the specimen well (S).
- 8) Observe for development of purple colored lines in the results window. Interpret test results at 15 minutes after adding assay buffer to the specimen well (S).

9) Do not interpret the test result after 25 minutes.



- Hold the specimen transfer device and assay buffer bottle vertically, else it can lead to inaccurate results
- Exactly 2 drops of assay buffer should be added. Adding more than 2 drops of assay buffer may cause over flooding or reverse migration phenomenon, which may lead to inaccurate results of the test.
- · Adding less than 2 drops of assay buffer may cause improper migration and poor background clearance which may lead to inaccurate results of the test.
- Do not read the test result after 25 minutes. Reading the result after the 25 minutes may give inaccurate results. After recording the results, dispose of the used test device as biohazard waste.

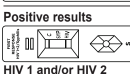
Internal Quality Control

The visualization of the purple colored Control Line in First Response® HIV 1+2 / Syphilis Combo Card Test indicates that the active ingredient of the strips are functional and the migration is successful. The control line is a procedural control serves to demonstrate functional reagents and correct migration of fluid.

How to Interpret test results

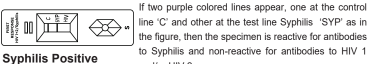


If only a single purple colored line appears, at control line "C" as in the figure, then the specimen is non-reactive for antibodies to Syphilis and HIV.

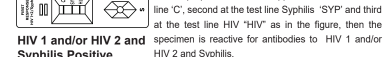


HIST CYSPONSE

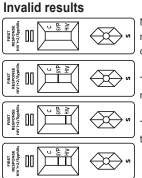
If two purple colored lines appear, one at the control line 'C' and other at the test line HIV "HIV" as in the figure, then the specimen is reactive for antibodies to HIV 1 and/or HIV 2 and non-reactive for antibodies to Syphilis



the figure, then the specimen is reactive for antibodies to Syphilis and non-reactive for antibodies to HIV 1 and/or HIV 2.



Syphilis Positive



No presence of purple colored control line 'C' in the results window (irrespective of the presence of purple colored test lines) indicates an invalid result.

Note: Interprete faint lines as the reactive lines.

The directions may not be followed correctly or the test may have deteriorated.

The Invalid test results should be retested with a new test device

Performance Characteristics

First Response® HIV 1+2 / Syphilis Combo Card Test has been tested using an in-house panel of Positive and Negative clinical specimens characterized by a commercial anti-HIV 1&2 ELISA kit and TPHA kit. First Response® HIV 1+2 / Syphilis Combo Card Test showed 100% sensitivity and 100% specificity. First Response® HIV 1+2 / Syphilis Combo Card Test showed 100% agreement with reference assays.

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po	On a simon dataile	First Response® HIV 1+2/Syphilis Combo Card						
Reference Method	Specimen details	HIV	HIV	Syphilis	Syphilis	Total		
æ-		Positive	Negative	Positive	Negative			
	51		· · ·					
	•	131	0	0	131	131		
		6	0	0	6	6		
		0	46	46	0	46		
	51	ositive Plas	ma specime					
		40	0	40	0	40		
	HIV and Syphilis Ne	egative Plas	ma specim	ens				
Ð	Negative Plasma Specimen	0	370	0	370	370		
labl	Total Plasma specimens	177	416	86	507	593		
avai	HIV Positive and Syphil	philis Negative Serum specimens						
all <u>y</u>	HIV 1 Positive Serum Specimen	419	0	0	419	419		
arci	HIV 2 Positive Serum Specimen	85	0	0	85	85		
Ĕ	Syphilis Positive and HIV Negative Serum specimens							
S	Syphilis Positive Serum Specimen	0	101	101	0	101		
DT	HIV and Syphilis N	HIV and Syphilis Negative Serum specimens						
AF	HIV Positive and Syphilis HIV 1 Positive Plasma Specimen HIV 2 Positive Plasma Specimen Syphilis Positive plasma Specimen HIV and Syphilis Positive and Syphilis Neg Negative Plasma Specimen Total Plasma specimens HIV 1 Positive Serum Specimen HIV 2 Positive Serum Specimen HIV 2 Positive Serum Specimen Syphilis Positive Serum Specimen Syphilis Positive Serum Specimen	0	3455	0	3455	3455		
LIS		504	3556	101	3959	4060		
ш	HIV Positive and Syphilis N	legative W	nole blood s	pecimens				
	HIV Positive Whole blood specimen	20	0	0	20	20		
	Syphilis Positive and H	IV Negative	Whole blo	od specime	ins			
		0	34	34	0	34		
	HIV and Syphilis Posit	tive Whole	olood speci	mens				
		31	0	31	0	31		
	HIV and Syphilis Nega	ative Whole	blood spec	imens				
		0	217	0	217	217		
	· ·	51	251	65	237	302		
-		•				002		

If three purple colored lines appear, one at the control line 'C', second at the test line Syphilis 'SYP' and third at the test line HIV "HIV" as in the figure, then the

Reference	Specimen details		First Response® HIV 1+2 / Syphilis Combo Card Test							
Method			Positive	Negative	Total	95% Confidence				
	Test Marker	Parameter	1 03/070	Negative	Result	Interval				
		Plasma Specimens								
ple	HIV	Sensitivity	177	00	177	(97.35%-100%)				
aila	TIIV	Specificity	00	416	416	(98.85%-100%)				
avi	Syphilis	Sensitivity	86	00	86	(94.67%-100%)				
Commercially available	Cypinio	Specificity	00	507	507	(99.06%-100%)				
erci	Serum Specimens									
E	HIV	Sensitivity	504	00	504	(99.05%-100%)				
Ō		Specificity	00	3556	3556	(99.86%-100%)				
10	Syphilis	Sensitivity	101	00	101	(95.43%-100%)				
ELISA/ RDT	Cypinio	Specificity	00	3959	3959	(99.87%-100%)				
ISI.		Whole blood	Specimens (C	apillary and v	enous bloo	d)				
	HIV	Sensitivity	51	00	51	(91.27%-100%)				
	1117	Specificity	00	251	251	(98.12%-100%)				
	Syphilis	Sensitivity	65	00	65	(93.04%-100%)				
	Syptillis	Specificity	00	237	237	(98.01%-100%)				

Seroconversion Panel Testing

The Analytical sensitivity of the First Response® HIV 1+2 / Syphilis Combo Card Test was carried out by testing commercially available Seroconversion panel. The commercially available HIV/Syphilis combo rapid lateral flow test was used as a reference kit for comparative performance study. Twenty-two (22) seroconversion panel was tested, in-house.

Analytical Sensitivity - In - House Evaluation								
Total Seroconversion Panels	Total		oonse® HIV 1+2 Combo Card Te		Reference HIV/Syphilis Combo rapid lateral flow test.			
	Specimens	Positive	Negative	Detection Index**	Positive	Negative	Detection Index**	
22	130	36	94	0.27	35	95	0.26	

** Detection Index = Total number of positive specimen by test kit / Total number of specimens.

Cross-Reactivity Study

First Response® HIV 1+2 / Syphilis Combo Card Test was tested with other diseases/conditions, which may give cross-reactivity with the test. The following 18 potential cross-reacting diseases/conditions did not affect the performance of the First Response[®] HIV 1+2 / Syphilis Combo Card Test.

Specimen details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive	Specimen details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive
P.falciparum Positive	05	Not Tested	05	Not Tested	HSV 1/2 Positive#	05	16	05	08
Pan Malaria Positive	05	Not Tested	05	Not Tested	HTLV- I Ab Positive#	07	08	07	04
Dengue NS1 Positive#	05	08	05	04	HTLV- II Ab Positive#	09	08	09	04
Pregnant Woman *	320	02	321	01	HSV - IIgG Positive#	08	08	08	04
CMV Positive#	03	08	03	04	Rubella IgG & IgM Positive#	15	16	15	08
ANA Positive#	04	08	04	04	HBV Positive [#]	103	08	103	04
HAV Positive#	04	08	04	04	Chikungunya Positive#	Not tested	08	Not tested	04
EBV Positive#	02	08	02	04	Anti-malarial drug medication#	04	08	04	04
HCV Positive#	103	08	103	04	Anti-TB drug medication#	05	10	05	05

Note : ^ Naturally appeared HIV and Syphilis positive specimens.

* Spiked HIV and Syphilis positive specimens.

Potential interference substances

The First Response® HIV 1+2 / Syphilis Combo Card Test was tested with potential interfering substances. The following 08 potential interfering substances did not affect the performance of First Response® HIV 1+2 / Syphilis Combo Card Test. However, Haemolysed specimens and lipaemic specimens showed poor background clearance, hence not recommended for testing. Lipaemic specimens can be used for the testing after centrifugation. Such specimens must be centrifuged at 5000 g for 10 minutes and use the supernatants for testing.

Specimen Details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive	Specimen Details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive
Lipaemic specimen**,#	25	08	25	04	Low Hematocrit specimens	05	Not tested	05	Not tested
Icteric specimens#	05	08	05	04	Whole blood specimen in ACD anticoagulant	182	Not tested	182	Not tested
Haemolytic specimens**	05	Not tested	05	Not tested	RF Ab Positive#	09	08	09	04
High Hematocrit specimens	05	Not tested	05	Not tested	dsDNA Antibody Positive Plasma [#]	01	08	01	04

Potential interference Drug substances

The details of interfering drug molecules are mentioned in the following table. Each interfering drug molecule substances were spiked at the final concentration of 250µg/ml in HIV 1, HIV 2 and Syphilis, positive as well as negative specimens, respectively. No false positive or false negative results were observed with any of drug molecules when tested with First Response® HIV 1+2 / Syphilis Combo Card Test.

Diclofenac	Naproxen IP	Acetaminophen	Rifampicin	Ibuprofen					
Folic acid	Pantoprazole	Pyrazinamide	Metformin	Aspirin					
Ecosprin	Hydrochlorothiazide	Cholecalciferol	Isoniazid	Ampicillin Sodium salt					
Magnesium sulphate	Ascorbic Acid (Limec)	Ritonavir	Ferrous Ascorbate	Nevirapine					
Daruvir		Cyclobenzaprine Hydrochloride							

Precision

The precision of the First Response[®] HIV 1+2 / Syphilis Combo Card Test was determined by using the 21 different specimens containing different concentrations of antibodies in 5 different replicates with 3 different lots of test devices. Between-run and within-run precision were observed 100%.

External Evaluation Report

Place of Evaluation	Year	Sensit	ivity	Specificity		
	rear	Syphilis	HIV	Syphilis	HIV	
Zimbabwe (Plasma)	2015	100% (92.94%-100%)	100% (95.60%-100%)	100% (98.00%-100%)	100% (97.59%-100%)	
Ghana (Serum/Plasma)	2017	100% (94.29%-100%)	100% (94.29%-100%)	100% (96.88%-100%)	100% (96.88%-100%)	
WHO evaluation (Serum/Plasma)	2018	99.0% (96.4% - 99.9%)	100% (98.2% - 100%)	100% (98.2% - 100%)	99.5% (97.2% - 100%	
Ghana (Capillary vs Venus whole blood specimen)	2018	100% (87.35%-100%)	100% (96.19%-100%)	100% (97.71%-100%)	100% (96.07%-100%	
Zimbabwe (Pregnant women whole blood specimen)	2019	100% (87.01%-100%)	100% (96.55%-100%)	100% (98.06%-100%)	100% (96.69%-100%	

Limitations

- 1) Do not use anti-coagulants other than heparin, EDTA, and sodium citrate.
- 2) Do not use the haemolysed specimen. A haemolysed specimen may give reddish background even after the end of test time.
- 3) Interpret a faint line as a positive line. Repeat the test in case of a very faint test line or if have any doubt for the test line.
- 4) Although a positive result may indicate an infection of HIV 1 and/or HIV 2 or Syphilis (Treponema pallidum), a diagnosis of diseases can only be made on clinical grounds. This test should not be used as the sole criteria for the diagnosis of HIV/ Treponema pallidum.
- 5) For confirmation, further analysis of the specimens should be performed, such as ELISA, or western blot analysis for HIV and TPHA for Syphilis. As with all diagnostic tests, results must be interpreted together with other clinical information available to the physician.
- 6) False negative results may arise because of hook effect due to a very high titer of antibody in a specimen. Repeat the test by using 1:10 dilution of the same specimen (01 portion) in respective non-reactive specimen matrix (09 portions)
- 7) A non-reactive result does not eliminate the possibility of infection with HIV1/2 and/or Treponema pallidum. The specimen may contain a low level of antibodies that cannot be detected by First Response® HIV 1+2 / Syphilis Combo Card Test. If a test result is non-reactive and clinical symptoms persists, additional testing using other reference method is recommended and/or retested for HIV antibodies after more than 21 days since the original testing.
- 8) Some HIV infected persons on antiretroviral medication may produce false negative results when tested with rapid diagnostic tests

SYMBOL LEGENDS

	Symbol	Explanation of symbol	Symbol	Explanation of symbol
	symbol		Symbol	. ,
	[]i	Consult instructions for use	₹	Contains sufficient for < n > tests
	NON	Non Sterile	REF	Product Code
	IVD	In vitro diagnostic medical device	LOT	Lot Number
4°(c 30°C	Store at 4-30 °C	444	Manufacturer
	\bigcirc	Caution	M	Date of manufacture (YYYY-MM)
	Ч,	Keep dry		Expiration Date (YYYY-MM)
	\otimes	Do not reuse	8	Do not use if test device pouch is damaged
	¥.	Keep away from sunlight		

References:

- 1) Hook EW et al. 2002. A randomized, comparative pilot study of azithro mycin versus benzathine penicillin G for treatment of early Syphilis. Sexually Transmitted Diseases. 8: 486-90.
- 2) Universal Access Report, Scaling up priority HIV/AIDS interventions in the health sector, Progress report 2010.
- 3) UNAIDS, 2013. Report on the global AIDS epidemic "GLOBAL REPORT'
- 4) Kieffer M. 2005. Mortality of infants born to HIV-infected mothers in Africa. The Lancet, 365(9454):120-121
- 5) WHO, 2007. The global elimination of congenital Syphilis: rationale and strategy for action
- 6) WHO, 2011. Sexually transmitted infections. Geneva: World Health Organization.
- 7) Aledort JE et al. 2006. Reducing the burden of sexually transmitted infections in resource-limited settings: the role of improved diagnostics. Nature, 444: 59-72.
- 8) Peeling RW, 2009. Utilization of rapid tests for sexually transmitted infections: promises and challenges. Infectious Diseases Journal, 3: 156-163.
- 9) Newcombe, Robert G. "Two-Sided Confidence Intervals for the Single Proportion: Comparison of Seven Methods," Statistics in Medicine, 17, 857-872 (1998)
- 10) Wilson, E. B. "Probable Inference, the Law of Succession, and Statistical Inference," Journal of the American Statistical Association 22 209-212 (1927)
- 11) TGS-5: Designing Instruction for use for in vitro diagnostic medical devices
- 12) A Short guide on methods: Measuring the impact of national PMTCT programmes (2012 Julv).

13) http://vassarstats.net/clin1.html#def , Richard Lowry.

14) Mwumvaneza Mutagoma, Eric Remera, Dieudonné Sebuhoro, Steve Kanters, David J. Riedel, and Sabin Nsanzimana, "The Prevalence of Syphilis Infection and Its Associated Factors in the General Population of Rwanda: A National Household-Based Survey," Journal of Sexually Transmitted Diseases, vol. 2016, Article ID 4980417, 8 pages, 2016. https://doi.org/10.1155/2016/4980417.

Product Disclaimer & Warnings

Every warnings and precaution should be taken into consideration before using the test. Failure to consider "Precaution, Warning, and Limitations" may not ensure the diagnostic ability and accuracy of this product. The test result may accordingly be affected by environmental factors and/or user error outside of the control of the Manufacturer and Distributor.

A definitive clinical diagnosis should not be based on the results of a single test, but it should be made by the physician after all clinical and laboratory findings have been evaluated

"In no event shall our company or its distributor is liable for any direct, indirect, punitive, unforeseen, incidental, special consequential damages, to property or life, whatsoever arising out of or connected with an incorrect diagnosis, whether positive or negative, in the use or misuse of this product". In the event of performance changes or product malfunction, please contact manufacturer.



· ISO 13485 & EN ISO 13485 Certified Company

ENGLISH Part No.(S)I20-INS-007. Rev.: AB. Date:2020-02-17 Note : Instructions for use will be printed in local language of the country using the test, if required.

FIRST RESPONSE® HIV 1+2 / SYPHILIS COMBO CARD TEST Rapid immunochromatographic Card Test for detection of Antibodies to HIV and/or Syphilis in human whole blood/ serum/ plasma

REF 120FRC25, 120FRC30, 120FRC50, 120FRC60 & 120FRC100

Intended Use

First Response® HIV 1+2 / Syphilis Combo Card Test is intended for us healthcare professionals and qualified laboratory personnel. It is a rapid, quality screening, in vitro diagnostic test for the detection of antibodies (IgG & IgM) spe to HIV (type 1 & 2) and Treponema pallidum in human serum, plasma or ven and capillary whole blood. The test can be used as an aid in the diagnosis of and/or Syphilis. The product can be used for symptomatic, asymptomatic pregnant women population. The test kit is not automated and does not require additional instrument. Reactive specimens should be confirmed by suppleme testing with ELISA, Western Blot or TPHA.

Introduction

HIV (Human Immunodeficiency Virus) is recognized as the etiologic age Acquired Immune Deficiency Syndrome (AIDS). The virus is transmitted by se contact, exposure to infected blood, certain body fluids or tissues, and from me to fetus or child during the perinatal period.

Syphilis is a venereal disease caused by the spirochete bacterium Trepon pallidum. It is ordinarily transmitted by sexual contact. It can also be transm congenitally by the transplacental passage of mother to the fetus and by b transfusion. In a case where a patient is infected with HIV as well as Syphi increases the chances of HIV transmission by increasing viral shedding seminal viral load. The prevalence of HIV is 3 times more in patients infected Syphilis compared to those not infected with Syphilis infection(14). Incorpor Syphilis screening in HIV prevention programs will help to prevent mother to transmission of HIV and Syphilis. This can be achieved by the implementation simple and affordable dual testing strategy for HIV and Syphilis which could imp screening uptake and accessibility of testing to accelerate time to treatment. WHO has reported a significantly high number of HIV and Syphilis co-infecti mother to child transmission (MTCT) in Africa Therefore, the WHO has annou in June 2012 that Prevention of Mother to Child Transmission (PMTCT) shoul be considered alone for HIV but considered for HIV and/or Syphilis both, w vision to eliminate new HIV infections to children by 2015(12). To achieve this v each pregnant woman should be tested for Syphilis and HIV both rather than only. Development of a single test device containing HIV and Syphilis antigen solve the issue defined above and will also be a useful step in achieving W ambitious goal.

Assay Principle

First Response® HIV 1+2 / Syphilis Combo Card Test is based on the princip immunochromatography for the qualitative detection of antibodies(IgG & specific for HIV 1&2 and/or Syphilis. The nitrocellulose membrane is coated w cocktail of recombinant antigen for HIV 1 (gp41) and HIV 2 (gp36) at test line and Recombinant TP antigen (P47, P45, P17, P15) specific for Treponema pal at the test line "Syp" and control reagent coated at the control line "C". When se or plasma or whole blood specimen is applied to the specimen well of the device, the cocktail of recombinant HIV 1+2 (gp41 & gp36) antigen - colloidal conjugate (CGC) & recombinant Treponema pallidum antigens colloidal conjugate will react with HIV and/or Syphilis specific antibodies, if present in specimen. The antibody-CGC antigen complex and assay buffer move along membrane chromatographically to the test regions and form a visible purple col line as the antigen-antibody-CGC antigen complex forms with a high degr sensitivity and specificity. If the specimen contains antibodies to Trepor pallidum, the purple colored line will appear in the test area at test line ' corresponding to the Syphilis line. If the specimen contains antibodies to H and/or 2, the purple colored line will appear in the test area at test line corresponding to HIV 1+2 line.

The presence of both test lines indicates that the specimen contains antibodi HIV as well as Treponema pallidum. The absence of the purple colored line at test line regions indicates that the specimen is non-reactive for HIV and Trepon pallidum, showing a negative result. The purple colored Control line will ap irrespective of a reactive or non-reactive specimen. The control line is a proceed control, serves to demonstrate functional reagents and correct migration of fluid.



N	laterials Provide	d				
		Washing!	PIBST RESPONSE HIV 1+2/SypNils		Alcohol Swab	FRST RESPONSE" HV 1+2 / SYTHULS For detailer of following to the web parts processing corrects, corrects, corrects protection
	HIV 1+2 / SYPHILIS Combo Card Test Anti HIV 1+2 / SYPHILIS Antibody Card Test Whole Biode (Seruer) Plasma	and the second			isopropyl Alcohol	International and the second secon
		Water States		Assay	1111	replanatio, appropriate and program server population. The te- submeted and data to replace any articular instances. It also been shared to confirm the population of and only and 10.000 to a or 1940. A shared because of the state of the st
	Preview Medical Corporation Private Limited *********************************	And Designation		Buffer	For Disinfection Use	quart invesse laborary Ryannes (alle). The sites is trave- net entert, approach is obtained inton, made heady fails a tra- mmer is basis within all of principal points. philo is a memorial channe massel by the particular points indees it is methodic traventioned by amount content in the indees of a methodic by the traventioned points of methods on particular by the traventioned by amount content in the mobiles of particular by the traventioned points of methods on particular size of the size o
	Test device pouch	Desiccant	As As	say buffer Alc	ohol Swab	It is the strategistics in the same strate a patient is information for Septimics, is increased for a first transmission by increas- ting patient sectors into the strategistic of the strategistic and information of the patient patient from the information of the patient floating and a patient patient from the information of the patient floating and the patient patient patient patient of the strategistic of the strategistic of the strategistic of states on information and on the strategistic patient patient information of the strategistic of
α				bottle		Instructions
	Specimen	-	<u>s</u>	Sterile	Twist Lancet	for use
	transfer device		Test device			
No	te: Materials provided othe	r than assay b	ouffer bottle ar	e for single use	only.	
Ma	aterials provided	I20FRC25	I20FRC30	120FRC50	120FRC60	I20FRC10
	st device pouch containing: est device, 1 desiccant	25 Nos.	30 Nos.	50 Nos.	60 Nos.	100 Nos.
	ecimen transfer device	25 Nos.	30 Nos.	50 Nos.	60 Nos.	100 Nos.
<u> </u>	say buffer bottle (2.5 ml)	1 No.	1 No.	2 Nos.	4 Nos.	4 Nos.
	erile twist lancets	25 Nos.	30 Nos.	50 Nos.		100 Nos.
	cohol swabs	25 Nos. 25 Nos.	30 Nos. 30 Nos.		60 Nos. 60 Nos.	100 Nos.
	tructions for use	25 NOS. 1 No.	1 No.	50 Nos. 1 No.	1 No.	2 Nos.
		-	-		1 110.	2 NUS.
N	laterials Require	d but No	ot Provide	ed		
•	New pair of disposab	le gloves a	nd face ma	sk for each t	est conducte	ed/specime
	collected by Fingerst	ick.				
•	Sterile gauze pad an	d tissue pa	iper.			
•	Permanent marker p	en and time	er.			
•	Extra sterile twist la	ncets, alco	ohol swabs	, and specir	nen transfe	r devices,
	needed.					
•	Sharp disposable bo					
•	Venipuncture blood o		it (if whole I	blood is colle	cted by ven	ipuncture)
S	torage and Stabi	-				
1)	First Response [®] HI	V 1+2 / Sy	philis Coml	oo Card Tes	t kit should	be stored
	4-30°C.					
2)	Do not freeze the ki					
3)	The kit is sensitive t	•		00 not store t	he kit at the	temperatu
	above 30°C and in I					
4)	Assay buffer (opene	-	,	-		e stable u
- ·	the expiry date print					
5)	Perform the test imr			0		
	pouch. If the desicc	ant color ha	as changed	trom orange	to green, d	o not use t
0.	test device.					
6)	Test device is sta		he printed	expiry date	on the po	ouch/extern
	secondary packagir	ng.				
	recautions					
	Wear protective glow			-	-	
	Dispose of used glov			e. Wash har	nds thorough	nly afterwa
	Avoid splashing or a					
	Clean up spills thoro					
5)	Decontaminate and	-	-			
	and specimen trans					
	container. Dispose o	f used ster	ile twist land	cets in a shar	ps box and	face mask
	a waste container.					
V	Varnings					
1)	For in vitro diagnos	-				
2)	Read the instructio	ns carefull	y before pe	erforming the	e test, any o	deviation v
,	invalidate the test r	esults.				
ý 3)	invalidate the test n Apply standard bio		cautions for	handling an	d disposal o	of potentia
,		safety prec		-	-	-
,	Apply standard bio	safety prec		-	-	-
,	Apply standard bios infective materials	safety prec including h	numan biol	ogical specir	nens irrespe	ective of t
3)	Apply standard bio infective materials disease state.	safety prec including h say buffer. I	numan biol It contains s	ogical specir sodium azide	nens irrespo as a preser	ective of the
3)	Apply standard bio infective materials disease state. Do not drink the ass	safety prec including h say buffer. I	numan biol It contains s	ogical specir sodium azide	nens irrespo as a preser	ective of t

5) Devices and assay buffer of a different lot must not be used.

- 6) Do not use the test device if the pouch is not intact.
- 7) Do not use the auto safety lancet if the lancet found uncapped (Refer specimen collection section)
- 8) Do not use the test device if the desiccant color has changed from orange to green.
- 9) Do not smoke, eat or drink while handling specimens and performing a test.
- 10) Do not re-use the test device, alcohol swab, auto safety lancet, and specimen transfer device as these are intended for single use only.
- 11) Perform the test by using kit assay buffer, any other buffer or fluid will invalidate the test results
- 12) Do not allow the tip of assay buffer bottle to touch specimen well as it may contaminate the assav buffer.
- 13) Do not use the test device and assay buffer beyond the date of expiry.
- 14) Do not eat the desiccant.
- 15) Do not use any other specimen other than human Whole blood/Serum/Plasma. Do not mix and interchange different specimens.

Specimen Collection

- 1) Venous blood collection: Collect the Whole blood in the collection tubes containing anticoagulants like EDTA, Heparin, Sodium citrate or ACD by venipuncture
- 2) Plasma collection: Collect the Whole blood in the collection tubes containing anticoagulants like EDTA, Heparin, Sodium citrate or ACD by venipuncture and centrifuge it at 3000 g for 10-15 minutes to obtain Plasma.
- 3) Serum collection: Collect Whole blood in the collection tubes without having any anticoagulants by venipuncture. Keep it in standing position for 30 minutes and centrifuge it at 3000 g for 10-15 minutes to obtain serum.

4) Capillary whole blood specimen collection:

- Wear gloves and massage the fingertip gently. It will help to obtain a round drop of blood.
- Wipe the complete fingertip with the alcohol swab provided and wait until the fingertip dried completely.

Auto Safety Lancet (Sterile Pressure Activated Lancet)

3 Dispose of safety lan in sharps Instructions for use \longrightarrow he clea against the

•Do not use the auto safety lancet if the auto safety lancet found uncapped. Detach the protective cap of the auto safety lancet provided. Squeeze the fingertip then push gently at the lateral side (avoid callus) of the fingertip as shown in above figure. Safely dispose of the used auto safety lancet in sharps container immediately after use.

•Wipe the first drop of the blood using sterile gauze. Without pressing too hard, gently squeeze fingertip once again to obtain second drop of blood(~40-50 µl).



• Take the specimen transfer device provided and hold it vertically. Gently squeeze the bulb of specimen transfer device and immerse open end in the center of a blood drop and release the bulb slowly to draw up the blood up to the 20 µl marking line on the specimen transfer device

Do not use the specimen transfer device having no marking. After completion of specimen collection, take the sterile gauze and apply pressure to the wound site to stop the bleeding. The specimen transfer device is for single use only

Note: Auto safety lancet is for single use only. Do not share used auto safety lancet with another person. Dispose of used auto safety lancet in sharp box and alcohol swab in biohazard waste container immediately after use.

Do not use expired auto safety lancet. Use of any expired auto safety lancet may cause infections at the punctured skin due to the expiry of its sterility. Use new auto safety lancet, alcohol swab and specimen transfer device and choose a different puncture site, if another finger pricking is required.

Specimen storage

1) Venous whole blood specimen should be used for testing immediately (within 1 hour) or shall be stored at 2-8°C for up to 72 hours (3 days). Do not use whole blood specimen stored for more than 3 days, it can cause a non-specific reaction. Do not freeze whole blood specimens.

Note: Mix the whole blood specimens in the tube by inverting the tube 3 or 4 times before use.

- 2) If serum or plasma specimens are not immediately tested, then they should be refrigerated at 2-8°C. For storage period greater than 72 hours (3 days), freezing at <-20°C is recommended up to 4 months.
- 3) Venous whole blood, serum and plasma specimens stored at 2-8 °C must be brought to room temperature before use. Serum or plasma specimens stored at ≤-20°C must be thawed at 15 to 25°C. Avoid more than 2 freeze-thaw cycles.

4) Serum or plasma specimens containing precipitate may yield inconsistent test results. Such specimens must be centrifuged at 5000 g for 10 minutes and then use clear supernatants for testing.

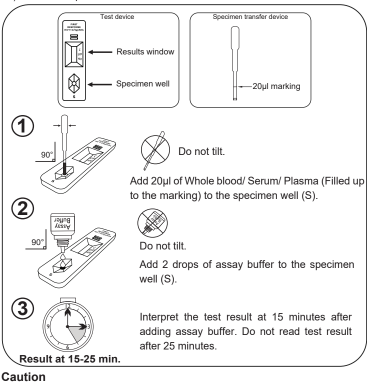
Test Procedure

- 1) Ensure that the test device & other components are at room temperature (15°C to 30°C) before starting the procedure.
- 2) Open the device pouch, take out the test device from aluminum pouch. Do not use the test device if the desiccant color has changed from orange to green.
- 3) Label the test device with the patient identification number. Place the test device on a flat, clean and dry surface.
- 4) Take out the specimen transfer device from plastic bag provided inside the kit. Gently squeeze the bulb of specimen transfer device and immerse the open end in the specimen and release the bulb slowly to draw up the serum/plasma/ capillary or venous whole blood up to 20µl marking line on the specimen transfer device.
- 5) Gently wipe away the excess specimen from the outer surface of the specimen transfer device with tissue paper before dispensing the specimen into the specimen well
- 6) Gently squeeze the bulb of specimen transfer device to add 20 µl of venous or capillary whole blood/ serum/ plasma to the specimen well by gently touching the tips of the specimen transfer device to the sample pad.

Caution: Dispose of used specimen transfer device and tissue paper as biohazard waste immediately after use

- 7) Hold the assay buffer bottle vertically and add two drops of assay buffer to the specimen well (S).
- 8) Observe for development of purple colored lines in the results window. Interpret test results at 15 minutes after adding assay buffer to the specimen well (S).

9) Do not interpret the test result after 25 minutes.



- · Hold the specimen transfer device and assay buffer bottle vertically, else it can lead to inaccurate results.
- Exactly 2 drops of assay buffer should be added. Adding more than 2 drops of assay buffer may cause over flooding or reverse migration phenomenon, which may lead to inaccurate results of the test.
- · Adding less than 2 drops of assay buffer may cause improper migration and poor background clearance which may lead to inaccurate results of the test.
- Do not read the test result after 25 minutes. Reading the result after the 25 minutes may give inaccurate results. After recording the results, dispose of the used test device as biohazard waste.

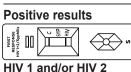
Internal Quality Control

The visualization of the purple colored Control Line in First Response® HIV 1+2 / Syphilis Combo Card Test indicates that the active ingredient of the strips are functional and the migration is successful. The control line is a procedural control serves to demonstrate functional reagents and correct migration of fluid.

How to Interpret test results

Negative results \Leftrightarrow

If only a single purple colored line appears, at control line "C" as in the figure, then the specimen is non-reactive for antibodies to Syphilis and HIV.

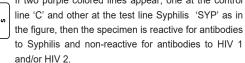


Syphilis Positive

Syphilis Positive

HIST CYSPONSE

line 'C' and other at the test line HIV "HIV" as in the figure, then the specimen is reactive for antibodies to HIV 1 and/or HIV 2 and non-reactive for antibodies to Syphilis If two purple colored lines appear, one at the control

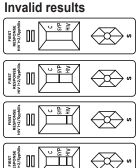


If three purple colored lines appear, one at the control \Leftrightarrow ÌŤŤ line 'C', second at the test line Syphilis 'SYP' and third

 \Leftrightarrow

at the test line HIV "HIV" as in the figure, then the HIV 1 and/or HIV 2 and specimen is reactive for antibodies to HIV 1 and/or HIV 2 and Svphilis

Note: Interprete faint lines as the reactive lines.



No presence of purple colored control line 'C' in the results window (irrespective of the presence of purple colored test lines) indicates an invalid result.

The directions may not be followed correctly or the test may have deteriorated.

The Invalid test results should be retested with a new test device

Performance Characteristics

First Response® HIV 1+2 / Syphilis Combo Card Test has been tested using an in-house panel of Positive and Negative clinical specimens characterized by a commercial anti-HIV 1&2 ELISA kit and TPHA kit. First Response® HIV 1+2 / Syphilis Combo Card Test showed 100% sensitivity and 100% specificity. First Response® HIV 1+2 / Syphilis Combo Card Test showed 100% agreement with reference assays.

	2									
od		First Res	sponse® HI\	/ 1+2/Syph	ilis Combo	Card				
Reference Method	Specimen details	HIV Positive	HIV Negative	Syphilis Positive	Syphilis Negative	Total				
Ľ.	HIV Positive and Syphil	hilis Negative Plasma specimens								
	HIV 1 Positive Plasma Specimen	131	0	0	131	131				
	HIV 2 Positive Plasma Specimen	6	0	0	6	6				
	Syphilis Positive and HI	V Negative	Plasma spe	ecimens	-					
	Syphilis Positive plasma Specimen	0	46	46	0	46				
	HIV and Syphilis P	ositive Plas	ma specime	ens						
	HIV and Syphilis Positive plasma Specimen	40	0	40	0	40				
	HIV and Syphilis Ne	egative Plas	sma specim	ens						
Ð	Negative Plasma Specimen	0	370	0	370	370				
ELISA/ RDT Commercially available	Total Plasma specimens	177	416	86	507	593				
ava	HIV Positive and Syphil	is Negative	Serum spe	cimens						
ally	HIV 1 Positive Serum Specimen	419	0	0	419	419				
erci	HIV 2 Positive Serum Specimen	85	0	0	85	85				
шш	Syphilis Positive and H	IV Negative	e Serum spe	ecimens	-	_				
ပိ	Syphilis Positive Serum Specimen	0	101	101	0	101				
RDT	HIV and Syphilis Negative Serum specimens									
SA/I	Negative Serum Specimen	0	3455	0	3455	3455				
ELIG	Total Serum specimens	504	3556	101	3959	4060				
	HIV Positive and Syphilis N		hole blood s	pecimens	-					
	HIV Positive Whole blood specimen	20	0	0	20	20				
	Syphilis Positive and H	-				-				
	Syphilis Positive Whole blood specimen	0	34	34	0	34				
	HIV and Syphilis Posi		blood speci	mens						
	HIV and Syphilis Positive Whole blood Specimen	31	0	31	0	31				
	HIV and Syphilis Neg	ative Whole		cimens						
	Negative Whole Blood Specimen	0	217	0	217	217				
	Total Whole blood specimens	51	251	65	237	302				

If two purple colored lines appear, one at the control

Reference	Specime	n details	First Res	onse [®] HIV 1	+2 / Syphili	s Combo Card Test				
Method	opecime		Positive	Negative	Total	95% Confidence				
	Test Marker	Parameter	1 001110	Nogativo	Result	Interval				
	Plasma Specimens									
ble	HIV	Sensitivity	177	00	177	(97.35%-100%)				
aila		Specificity	00	416	416	(98.85%-100%)				
avi	Syphilis	Sensitivity	86	00	86	(94.67%-100%)				
Commercially available	ojpinio	Specificity	00	507	507	(99.06%-100%)				
erci	Serum Specimens									
шш	HIV	Sensitivity	504	00	504	(99.05%-100%)				
Ō	TIIV	Specificity	00	3556	3556	(99.86%-100%)				
ELISA/ RDT	Syphilis	Sensitivity	101	00	101	(95.43%-100%)				
× Γ	ojpinio	Specificity	00	3959	3959	(99.87%-100%)				
IS/	Whole blood Specimens (Capillary and venous blood)									
E	HIV	Sensitivity	51	00	51	(91.27%-100%)				
	піх	Specificity	00	251	251	(98.12%-100%)				
	Syphilis	Sensitivity	65	00	65	(93.04%-100%)				
	Syprims	Specificity	00	237	237	(98.01%-100%)				

Seroconversion Panel Testing

The Analytical sensitivity of the First Response® HIV 1+2 / Syphilis Combo Card Test was carried out by testing commercially available Seroconversion panel. The commercially available HIV/Syphilis combo rapid lateral flow test was used as a reference kit for comparative performance study. Twenty-two (22) seroconversion panel was tested, in-house.

Analytical Sensitivity - In - House Evaluation									
Total Seroconversion Panels	Total		oonse® HIV 1+2 Combo Card Te		Reference HIV/Syphilis Combo rapid lateral flow test.				
	Specimens	Positive	Negative	Detection Index**	Positive	Negative	Detection Index**		
22	130	36	94	35	95	0.26			

** Detection Index = Total number of positive specimen by test kit / Total number of specimens.

Cross-Reactivity Study

First Response® HIV 1+2 / Syphilis Combo Card Test was tested with other diseases/conditions, which may give cross-reactivity with the test. The following 18 potential cross-reacting diseases/conditions did not affect the performance of the First Response[®] HIV 1+2 / Syphilis Combo Card Test.

Specimen details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive	Specimen details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive
P.falciparum Positive	05	Not Tested	05	Not Tested	HSV 1/2 Positive#	05	16	05	08
Pan Malaria Positive	05	Not Tested	05	Not Tested	HTLV- I Ab Positive#	07	08	07	04
Dengue NS1 Positive#	05	08	05	04	HTLV- II Ab Positive#	09	08	09	04
Pregnant Woman ^	320	02	321	01	HSV - IIgG Positive#	08	08	08	04
CMV Positive#	03	08	03	04	Rubella IgG & IgM Positive#	15	16	15	08
ANA Positive#	04	08	04	04	HBV Positive [#]	103	08	103	04
HAV Positive#	04	08	04	04	Chikungunya Positive#	Not tested	08	Not tested	04
EBV Positive#	02	08	02	04	Anti-malarial drug medication#	04	08	04	04
HCV Positive#	103	08	103	04	Anti-TB drug medication#	05	10	05	05

Note : ^ Naturally appeared HIV and Syphilis positive specimens.

* Spiked HIV and Syphilis positive specimens.

Potential interference substances

The First Response® HIV 1+2 / Syphilis Combo Card Test was tested with potential interfering substances. The following 08 potential interfering substances did not affect the performance of First Response® HIV 1+2 / Syphilis Combo Card Test. However, Haemolysed specimens and lipaemic specimens showed poor background clearance, hence not recommended for testing. Lipaemic specimens can be used for the testing after centrifugation. Such specimens must be centrifuged at 5000 g for 10 minutes and use the supernatants for testing.

Specimen Details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive	Specimen Details	HIV Negative	HIV Positive	Syphilis Negative	Syphilis Positive
Lipaemic specimen**,#	25	08	25	04	Low Hematocrit specimens	05	Not tested	05	Not tested
Icteric specimens#	05	08	05	04	Whole blood specimen in ACD anticoagulant	182	Not tested	182	Not tested
Haemolytic specimens**	05	Not tested	05	Not tested	RF Ab Positive#	09	08	09	04
High Hematocrit specimens	05	Not tested	05	Not tested	dsDNA Antibody Positive Plasma [#]	01	08	01	04

Potential interference Drug substances

The details of interfering drug molecules are mentioned in the following table. Each interfering drug molecule substances were spiked at the final concentration of 250µg/ml in HIV 1, HIV 2 and Syphilis, positive as well as negative specimens, respectively. No false positive or false negative results were observed with any of drug molecules when tested with First Response[®] HIV 1+2 / Syphilis Combo Card Test.

Diclofenac	Naproxen IP	Acetaminophen	Rifampicin	Ibuprofen					
Folic acid	Pantoprazole	Pyrazinamide	Metformin	Aspirin					
Ecosprin	Hydrochlorothiazide	Cholecalciferol	Isoniazid	Ampicillin Sodium salt					
Magnesium sulphate	Ascorbic Acid (Limec)	Ritonavir	Ferrous Ascorbate	Nevirapine					
Daruvir		Cyclobenzaprine Hydrochloride							

Precision

The precision of the First Response[®] HIV 1+2 / Syphilis Combo Card Test was determined by using the 21 different specimens containing different concentrations of antibodies in 5 different replicates with 3 different lots of test devices. Between-run and within-run precision were observed 100%.

External Evaluation Report

Place of Evaluation	Year	Sensit	ivity	Specificity		
	rour	Syphilis	HIV	Syphilis	HIV	
Zimbabwe (Plasma)	2015	100% (92.94%-100%)	100% (95.60%-100%)	100% (98.00%-100%)	100% (97.59%-100%)	
Ghana (Serum/Plasma)	2017	100% (94.29%-100%)	100% (94.29%-100%)	100% (96.88%-100%)	100% (96.88%-100%)	
WHO evaluation (Serum/Plasma)	2018	99.0% (96.4% - 99.9%)	100% (98.2% - 100%)	100% (98.2% - 100%)	99.5% (97.2% - 100%	
Ghana (Capillary vs Venus whole blood specimen)	2018	100% (87.35%-100%)	100% (96.19%-100%)	100% (97.71%-100%)	100% (96.07%-100%	
Zimbabwe (Pregnant women whole blood specimen)	2019	100% (87.01%-100%)	100% (96.55%-100%)	100% (98.06%-100%)	100% (96.69%-100%	

Limitations

- 1) Do not use anti-coagulants other than heparin, EDTA, and sodium citrate.
- 2) Do not use the haemolysed specimen. A haemolysed specimen may give reddish background even after the end of test time.
- Interpret a faint line as a positive line. Repeat the test in case of a very faint test line or if have any doubt for the test line.
- 4) Although a positive result may indicate an infection of HIV 1 and/or HIV 2 or Syphilis (*Treponema pallidum*), a diagnosis of diseases can only be made on clinical grounds. This test should not be used as the sole criteria for the diagnosis of HIV/ *Treponema pallidum*.
- 5) For confirmation, further analysis of the specimens should be performed, such as ELISA, or western blot analysis for HIV and TPHA for Syphilis. As with all diagnostic tests, results must be interpreted together with other clinical information available to the physician.
- 6) False negative results may arise because of hook effect due to a very high titer of antibody in a specimen. Repeat the test by using 1:10 dilution of the same specimen (01 portion) in respective non-reactive specimen matrix (09 portions).
- 7) A non-reactive result does not eliminate the possibility of infection with HIV1/2 and/or *Treponema pallidum*. The specimen may contain a low level of antibodies that cannot be detected by First Response[®] HIV 1+2 / Syphilis Combo Card Test. If a test result is non-reactive and clinical symptoms persists, additional testing using other reference method is recommended and/or retested for HIV antibodies after more than 21 days since the original testing.
- Some HIV infected persons on antiretroviral medication may produce false negative results when tested with rapid diagnostic tests.

SYMBOL LEGENDS

Symbol Explanation of symbol Symbol Explanation of symbol									
Explanation of symbol	Symbol	Explanation of symbol							
Consult instructions for use	₹ T	Contains sufficient for < n > tests							
Non Sterile	REF	Product Code							
In vitro diagnostic medical device	LOT	Lot Number							
Store at 4-30 °C	444	Manufacturer							
Caution	~~	Date of manufacture (YYYY-MM)							
Keep dry		Expiration Date (YYYY-MM)							
Do not reuse		Do not use if test device pouch is damaged							
Keep away from sunlight									
	Non Sterile In vitro diagnostic medical device Store at 4-30 °C Caution Keep dry Do not reuse	Consult instructions for use Image: Consult instructions for use Non Sterile Image: Consult instructions for use In vitro diagnostic medical device Image: Consult instruction Store at 4-30 °C Image: Consult instruction Caution Image: Consult instruction Keep dry Image: Consult instruction Do not reuse Image: Consult instruction							

References:

- Hook EW et al. 2002. A randomized, comparative pilot study of azithro mycin versus benzathine penicillin G for treatment of early Syphilis. Sexually Transmitted Diseases. 8: 486-90.
- Universal Access Report, Scaling up priority HIV/AIDS interventions in the health sector, Progress report 2010.
- 3) UNAIDS, 2013. Report on the global AIDS epidemic "GLOBAL REPORT"
- Kieffer M. 2005. Mortality of infants born to HIV-infected mothers in Africa. The Lancet, 365(9454):120-121.
- WHO, 2007. The global elimination of congenital Syphilis: rationale and strategy for action.
- 6) WHO, 2011. Sexually transmitted infections. Geneva: World Health Organization.
- Aledort JE et al. 2006. Reducing the burden of sexually transmitted infections in resource-limited settings: the role of improved diagnostics. Nature, 444: 59-72.
- Peeling RW, 2009. Utilization of rapid tests for sexually transmitted infections: promises and challenges. Infectious Diseases Journal, 3: 156-163.
- Newcombe, Robert G. "Two-Sided Confidence Intervals for the Single Proportion: Comparison of Seven Methods," Statistics in Medicine, 17, 857-872 (1998).
- Wilson, E. B. "Probable Inference, the Law of Succession, and Statistical Inference," Journal of the American Statistical Association, 22, 209-212 (1927).
- 11) TGS-5: Designing Instruction for use for in vitro diagnostic medical devices.
- A Short guide on methods: Measuring the impact of national PMTCT programmes (2012 July).

13) http://vassarstats.net/clin1.html#def , Richard Lowry.

14) Mwumvaneza Mutagoma, Eric Remera, Dieudonné Sebuhoro, Steve Kanters, David J. Riedel, and Sabin Nsanzimana, "The Prevalence of Syphilis Infection and Its Associated Factors in the General Population of Rwanda: A National Household-Based Survey," Journal of Sexually Transmitted Diseases, vol. 2016, Article ID 4980417, 8 pages, 2016. https://doi.org/10.1155/2016/4980417.

Product Disclaimer & Warnings

Every warnings and precaution should be taken into consideration before using the test. Failure to consider "Precaution, Warning, and Limitations" may not ensure the diagnostic ability and accuracy of this product. The test result may accordingly be affected by environmental factors and/or user error outside of the control of the Manufacturer and Distributor.

A definitive clinical diagnosis should not be based on the results of a single test, but it should be made by the physician after all clinical and laboratory findings have been evaluated.

"In no event shall our company or its distributor is liable for any direct, indirect, punitive, unforeseen, incidental, special consequential damages, to property or life, whatsoever arising out of or connected with an incorrect diagnosis, whether positive or negative, in the use or misuse of this product". In the event of performance changes or product malfunction, please contact manufacturer.



ISO 13485 & EN ISO 13485 Certified Company

Part No.(S)I20-INS-011, Rev.: AB, Date:2021-12-15 ENGLISH Note : Instructions for use will be printed in local language of the country using the test, if required.

FIRST RESPONSE® HIV 1+2 / SYPHILIS COMBO CARD TEST

Rapid immunochromatographic Card Test for detection of Antibodies to HIV and/or Syphilis in human whole blood/ serum/ plasma

 REF
 I20FRC25-SA

 Image: Second Se

Intended Use

First Response® HIV 1+2 / Syphilis Combo Card Test is intended for use the healthcare professionals and qualified laboratory personnel. It is a rapid, qualitative screening, in vitro diagnostic test for the detection of antibodies (IgG & IgM) specified to HIV (type 1 & 2) and *Treponema pallidum* in human serum, plasma or venous and capillary whole blood. The test can be used as an aid in the diagnosis of H and/or Syphilis. The product can be used for symptomatic, asymptomatic arpregnant women population. The test kit is not automated and does not require an additional instrument. Reactive specimens should be confirmed by supplement testing with ELISA, Western Blot or TPHA.

Introduction

HIV (Human Immunodeficiency Virus) is recognized as the etiologic agent Acquired Immune Deficiency Syndrome (AIDS). The virus is transmitted by sex contact, exposure to infected blood, certain body fluids or tissues, and from mot to fetus or child during the perinatal period.

Syphilis is a venereal disease caused by the spirochete bacterium Trepone pallidum. It is ordinarily transmitted by sexual contact. It can also be transmit congenitally by the transplacental passage of mother to the fetus and by blo transfusion. In a case where a patient is infected with HIV as well as Syphilis increases the chances of HIV transmission by increasing viral shedding seminal viral load. The prevalence of HIV is 3 times more in patients infected Syphilis compared to those not infected with Syphilis infection(14). Incorporat Syphilis screening in HIV prevention programs will help to prevent mother to ch transmission of HIV and Syphilis. This can be achieved by the implementation simple and affordable dual testing strategy for HIV and Syphilis which could impre screening uptake and accessibility of testing to accelerate time to treatment. WHO has reported a significantly high number of HIV and Syphilis co-infectior mother to child transmission (MTCT) in Africa Therefore, the WHO has announ in June 2012 that Prevention of Mother to Child Transmission (PMTCT) should be considered alone for HIV but considered for HIV and/or Syphilis both, with vision to eliminate new HIV infections to children by 2015(12). To achieve this vis each pregnant woman should be tested for Syphilis and HIV both rather than only. Development of a single test device containing HIV and Syphilis antigens solve the issue defined above and will also be a useful step in achieving WHG ambitious goal.

Assay Principle

First Response® HIV 1+2 / Syphilis Combo Card Test is based on the principle immunochromatography for the qualitative detection of antibodies(IgG & Ig specific for HIV 1&2 and/or Syphilis. The nitrocellulose membrane is coated wit cocktail of recombinant antigen for HIV 1 (gp41) and HIV 2 (gp36) at test line "H and Recombinant TP antigen (P47, P45, P17, P15) specific for Treponema pallid at the test line "Syp" and control reagent coated at the control line "C". When ser or plasma or whole blood specimen is applied to the specimen well of the device, the cocktail of recombinant HIV 1+2 (gp41 & gp36) antigen - colloidal g conjugate (CGC) & recombinant Treponema pallidum antigens colloidal o conjugate will react with HIV and/or Syphilis specific antibodies, if present in specimen. The antibody-CGC antigen complex and assay buffer move along membrane chromatographically to the test regions and form a visible purple colo line as the antigen-antibody-CGC antigen complex forms with a high degree sensitivity and specificity. If the specimen contains antibodies to Trepone pallidum, the purple colored line will appear in the test area at test line "Sy corresponding to the Syphilis line. If the specimen contains antibodies to HIV and/or 2, the purple colored line will appear in the test area at test line "H corresponding to HIV 1+2 line.

The presence of both test lines indicates that the specimen contains antibodies HIV as well as *Treponema pallidum*. The absence of the purple colored line at bot test line regions indicates that the specimen is non-reactive for HIV and *Treponen pallidum*, showing a negative result. The purple colored Control line will apper irrespective of a reactive or non-reactive specimen. The control line is a procedu control, serves to demonstrate functional reagents and correct migration of fluid.



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