# WHO Prequalification of In Vitro Diagnostics PUBLIC REPORT

# Product: One Step HIV1/2 Whole Blood/Serum/Plasma Test WHO reference number: PQDx 0357-004-00

One Step HIV1/2 Whole Blood/Serum/Plasma Test with product codes W006-C4P2, W006-P0045, W006-P0046, W006-P0047, W006-P0048, W006-C4P2-F, W006-P0049, W006-P0050, W006-P0051 and W006-P0052, manufactured by Guangzhou Wondfo Biotech Co., Ltd, Rest-of-World regulatory version, was accepted for the WHO list of prequalified in vitro diagnostics and was listed on 29 November 2018.

# Summary of WHO prequalification assessment for One Step HIV1/2 Whole Blood/Serum/Plasma Test

	Date	Outcome
Prequalification listing	29-Nov- 2018	listed
Dossier assessment	26-Apr-2018	MR
Site inspection(s) of quality	12-Oct-2018	MR
management system		
Product performance	First quarter of 2018	MR
evaluation		

MR: Meets Requirements

# **Report amendments and/or product changes**

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	Addition of new product codes to One Step HIV1/2 Whole Blood/Serum/Plasma Test. Alcohol swab and safety lancet/ lancet will be included in the new product codes. Added product codes were, W006-P0045 (25 Tests/Kit), W006-P0046 (25 Tests/Kit), W006-P0047 (25 Tests/Kit), W006-P0048 (25 Tests/Kit), W006-P0049 (40 Tests/Kit), W006-P0050 (40 Tests/Kit), W006-P0051 (40 Tests/Kit) and W006- P0052 (40 Tests/Kit).	6-Jul-2020
3.0	Correction of product code from WP006-CP42 to W006-C4P2	26-Aug-2021

# Intended use:

According to the manufacturer's claim, "Wondfo One Step HIV 1/2 Whole Blood/Serum/Plasma Test is a manual qualitative in vitro diagnostic for the detection of HIV 1/2 antibodies in human venous whole blood, capillary whole blood, serum and plasma specimens. It is intended for professional use in either laboratory or point of care settings. It is intended for aiding the diagnosis of HIV infection in symptomatic, symptomatic populations and persons at risk of HIV infection. It is not intended for testing children below 2 years and it is not intended for blood donor screening".

# Assay description:

According to the manufacturer's claim, "Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test is a rapid immunochromatographic direct binding test for the visual detection of HIV antibodies in venous whole blood, fingerstick whole blood, serum or plasma samples in the diagnosis of HIV infection. Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test adopts double antigen sandwich method. When the specimen is added into the test device, the specimen is absorbed into the device by capillary action, mixes with the antigen-dye conjugate, and flows across the HIV antigen pre-coated membrane. When the HIV antibody levels are at or above the detection limit of the test, HIV antibodies in the specimen bind to the antigen-dye conjugate and are captured by antigen immobilized in the test region (T) of the device. This produces a colored test band and indicates a positive result. If no HIV antibodies are present or below the detection limit of the assay, no colored band will be visible in the test region (T) of the device. This indicates a negative result. To serve as a procedure control, a colored line will appear at the control region (C), if the test has been performed properly."

# Test kit contents

Components	Pouch	Buffer Instruction		Alcohol swab	Sterile lancet	Safety lancet (pcs)		
Catalog No.	( <b>pcs</b> )	(bottle)	( <b>pcs</b> )	( <b>pcs</b> )	( <b>pcs</b> )	18G	21G	23G
W006-C4P2(25 Tests/Kit)	25	1	1	-	-	-	-	-
W006-P0045 (25 Tests/Kit)	25	1	1	25	25	-	-	-
W006-P0046 (25 Tests/Kit)	25	1	1	25	-	25	-	-
W006-P0047 (25 Tests/Kit)	25	1	1	25	-	-	25	-
W006-P0048 (25 Tests/Kit)	25	1	1	25	-	-	-	25
W006-C4P2-F(40 Tests/Kit)	40	2	1	-	-	-	-	-
W006-P0049 (40 Tests/Kit)	40	2	1	40	40	-	-	-
W006-P0050 (40 Tests/Kit)	40	2	1	40	-	40	-	-
W006-P0051 (40 Tests/Kit)	40	2	1	40	-	-	40	-
W006-P0052 (40 Tests/Kit)	40	2	1	40	-	-	-	40

## Items required but not provided

- Specimen collection containers
- Centrifuge (for serum/plasma specimen only)
- Capillary tube (for fingerstick whole blood only)
- Timer
- Protective gloves
- Specimen and test waste container

# Storage

The test kit and sample diluent should be stored at 2 -30 °C.

# Shelf-life upon manufacture

24 months (test kit and buffer).

# Warnings/limitations

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

# Prioritization for prequalification

Based on the established eligibility criteria, One Step HIV1/2 Whole Blood/Serum/Plasma Test was given priority for WHO prequalification assessment.

# Dossier assessment

Guangzhou Wondfo Biotech Co., Ltd submitted a product dossier for One Step HIV1/2 Whole Blood/Serum/Plasma Test as per the "*Instructions for compilation of a product dossier*" (PQDx\_018 v1). 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 26 April 2018.

Based on the product dossier screening and assessment findings, the product dossier for One Step HIV1/2 Whole Blood/Serum/Plasma Test meets WHO prequalification requirements.

# Manufacturing site inspection

A comprehensive inspection was performed at the site(s) of manufacture (8 Lizhishan Road, Science City, Luogang District, Guangzhou, 510663, Republic of China) of **One Step HIV1/2 Whole Blood/Serum/Plasma Test** between 19-21 March 2018 as per the "Information for manufacturers on prequalification inspection procedures for the sites of manufacture of diagnostics" (PQDx\_014 v1). The inspection found that the manufacturer had an acceptable quality management system and good manufacturing practices in place that ensured the consistent manufacture of a product of good quality.

The manufacturer's responses to the nonconformities found at the time of the inspection were accepted on 12 October 2018.

Based on the site inspection and corrective action plan review, the quality management system for **One Step HIV1/2 Whole Blood/Serum/Plasma Test** meets WHO prequalification requirements.

# Product performance evaluation

One Step HIV 1/2 Whole Blood/Serum/Plasma Test (Guangzhou Wondfo Biotech Co., Ltd) was evaluated by WHO in the first quarter of 2018 using serum/plasma specimens. From this evaluation, we drew the following conclusions:

One Step HIV 1/2 Whole Blood/Serum/Plasma Test (Guangzhou Wondfo Biotech Co., Ltd) is a qualitative rapid immunochromatographic assay for the detection of HIV-1/2 antibodies in human serum/plasma and venous/capillary whole blood specimens.

A volume of 10  $\mu$ L of serum/plasma/whole blood is needed to perform the assay. This type of assay requires no sophisticated 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.

In this limited evaluation on a panel of 1199 clinically-derived specimens, we found:

Performance characteristics in comparison with an agreed reference standard						
	Initial (95% CI)	Final (95% CI)				
Sensitivity %	100.0% (99.2% - 100.0%)	100.0% (99.2% - 100.0%)				
Specificity %	100.0% (99.5% - 100.0%) 100.0% (99.5% - 10					
Invalid rate %	0%					
Inter-reader variability %	0%					

Lot to lot variation was acceptable for all specimens

Additional performance characteristics					
Sensitivity during seroconversion	Seroconversion sensitivity index of -0.125, therefore				
on eight seroconversion panels in	detection is 0.125 specimens earlier than the				
comparison with a benchmark	benchmark assay.				
assay; Enzygnost Anti-HIV 1/2 Plus					
(Siemens Healthcare Diagnostics)					
Analytical sensitivity on a mixed	5 of 6 specimens were correctly classified.				
titer panel in comparison with an					
agreed reference standard					
Lot to lot variation on a dilution	Acceptable				
panel in comparison with an					
agreed reference standard					

Key operational characteristics	
Validated specimen types	Serum, plasma (EDTA, heparin or sodium citrate), venous whole blood and capillary whole blood.
Number of steps	2 with precision required
Time to result	15 minutes

# Labelling

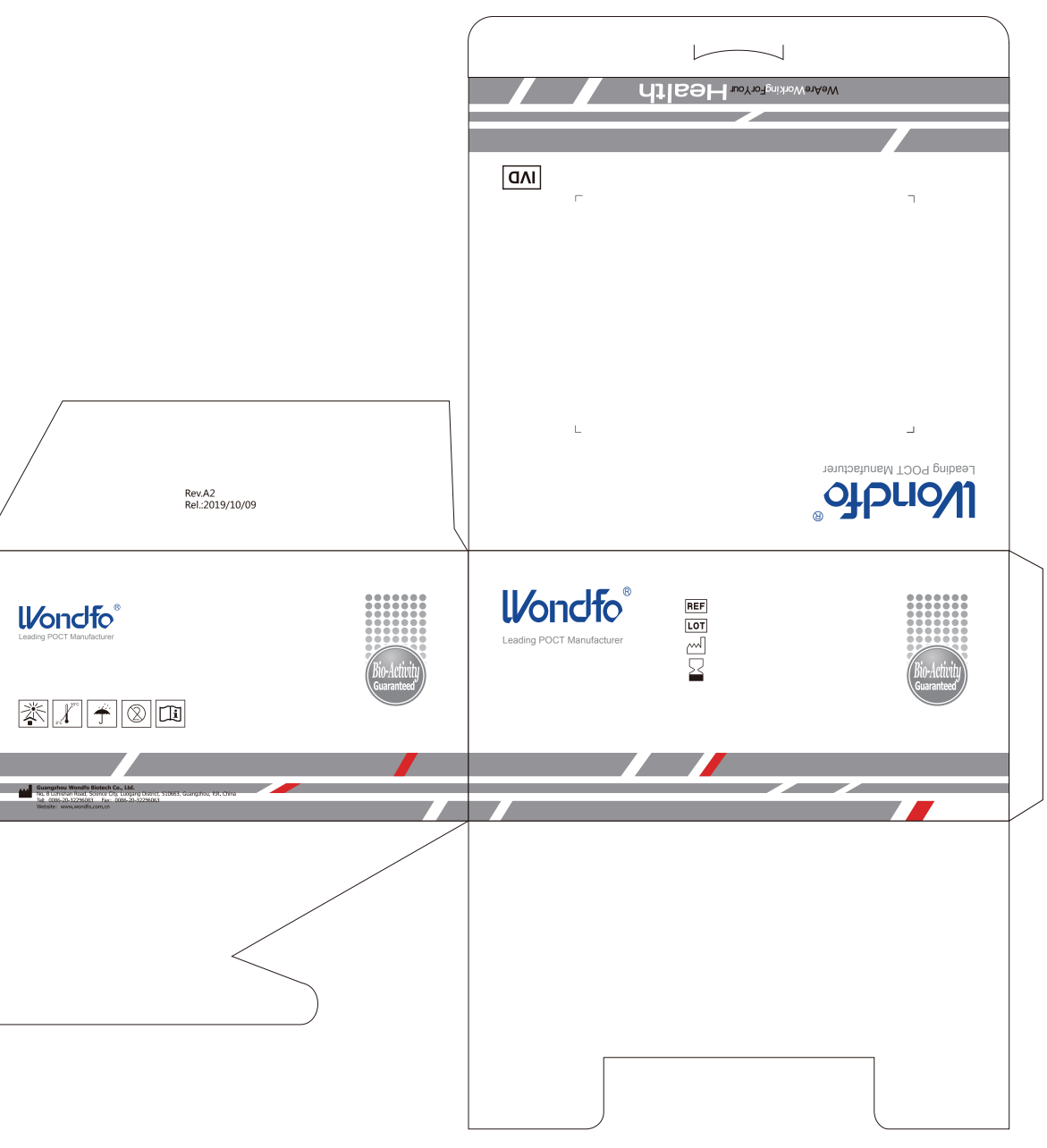
- 1. Labels
- 2. Instructions for use

1. Labels

1.1 Outside packaging box

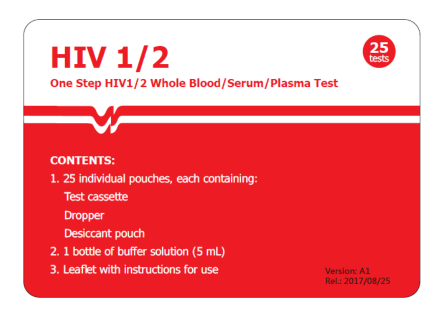
Leading POCT Manufacturer ll/ondfo<sup>®</sup> FOR IN VITRO DIAGNOSTIC USE ONLY FOR PROFESSIONAL USE ONLY Guangzhou Wondro Biotech Co., Ltd. No. 8 Lizhisharf Koad, Sciefice City, Luogang District, 510663, Guangzhou, P.R. Ci Tel: 0086-20-32295083 – Eax: 0086-20-32295063 Webcite, www.ewergite.com.org.

Size: 160\*140\*80mm

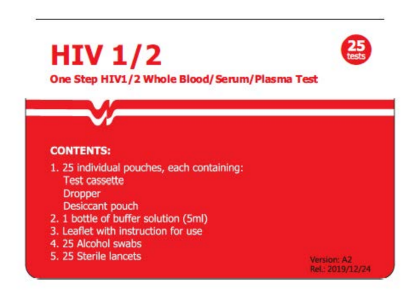


# **1.2** Contents label on outside packaging box

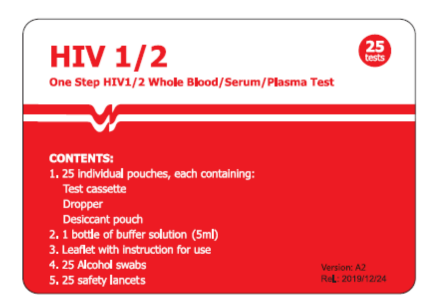
Product code W006-C4P2



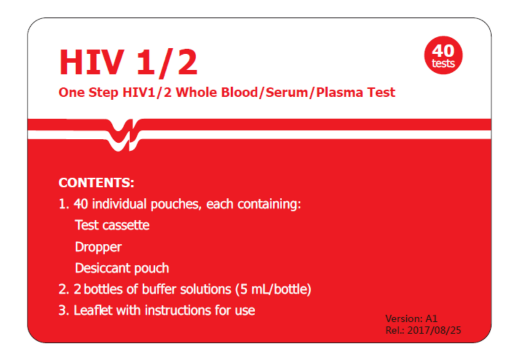
Product code W005-P0045



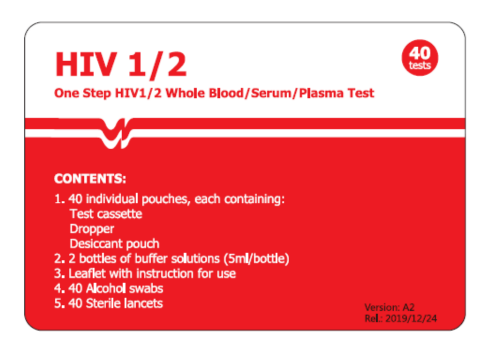
Product codes W006-P0046, W006-P0047 and W006-P0048



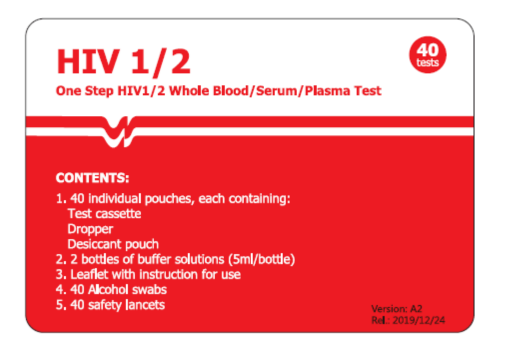
Product code W006-C4P2-F



# Product W006-P0049



Product codes W006-P0050, W006-P0051 and W006-P0052

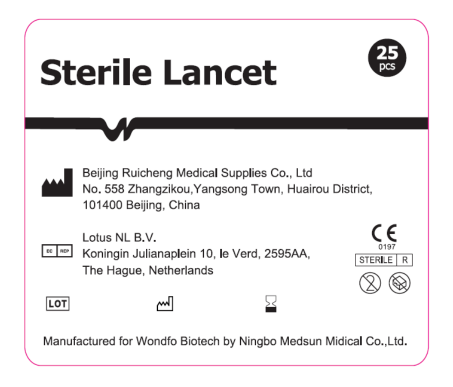


1.2 Buffer label

# Buffer for One Step HIV 1/2 Whole Blood/Serum/Plasma Test Lot: XXXXXXXX Exp: XXXX/XX/XX Vol: 5ml Store at: 2–30°C Version: A1 Rel: 2017/08/25

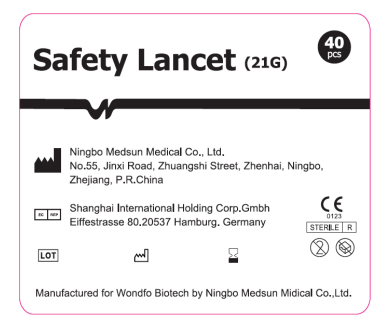
**1.3 Test device package** 

# 1.4 Sterile lancet label



# 1.5 Sterile safety lancets labels





# Product code W006-P0052

Safety Lancet (23G)							
•••	Ningbo Medsun Medical Co., Ltd. No.55, Jinxi Road, Zhuangshi Street, Zhenhai, Ningbo, Zhejiang, P.R.China						
EC REP	Shanghai International Holding Corp.Gmbh Eiffestrasse 80.20537 Hamburg. Germany						
LOT	₩ 🗳 🛞						
Manuf	actured for Wondfo Biotech by Ningbo Medsun Midical Co.,Ltd.						

# 1.5 Alcohol swab label

	cohol Swa Isopropyl Alcohol	b	<b>40</b> pcs			
<b>^</b>	SteriLance Medical (Suzhou) Inc No.68 Litanghe Road, Xiangchei		3 China			
EC REP	EMERGO EUROPE Prinsessegracht 20, 2514 AP, The Hague, The Netherlands		CE 0197 STERILE R			
LOT	ഷ	$\Sigma$	8 8			
Manufactured for Wondfo Biotech by Ningbo Medsun Midical Co.,Ltd.						

2.0 Instructions for use<sup>1</sup>

 $<sup>^1</sup>$  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.

# One Step HIV1/2 Wonclfo<sup>®</sup> Whole Blood/Serum/Plasma Test

#### Catalog No.

W006-C4P2 (25Tests) W006-C4P2-F(40tests) W006-P0045 (25Tests) W006-P0049 (40Tests) W006-P0046 (25Tests) W006-P0050 (40Tests) W006-P0047 (25Tests) W006-P0051 (40Tests) W006-P0048 (25Tests) W006-P0052 (40Tests)

#### INTENDED USE

Wondfo® One Step HIV 1/2 Whole Blood/Serum/Plasma Test is intended for use by trained users(in either laboratory or point-of-care settings), and is a qualitative, screening, in vitro diagnostic test for detection of HIV 1/2 antibodies in human venous whole blood, fingerstick whole blood, serum or plasma, aid to diagnosis of HIV infection.

#### SUMMARY AND EXPLANATION OF THE TEST

HIV (human immunodeficiency virus) is the pathogen of AIDS (acquired immunodeficiency syndrome). HIV belongs to Retroviridae genus Lentivirus family, and there are two groups of HIV, HIV-1 and HIV-2. HIV-1 is highly mutagenic and can be divided into 9 subtypes by the mutations in its membrane protein, which are A, B, C, D, E, F, G, H and O, HIV-2 has 60% nucleotide acid homology with HIV-1, but they are different in their ability of infection, HIV-1 is the most prevailing virus strain. Once infected, it mutates guickly and has bad prognosis. HIV-2 has a longer latent period, and relative weaker in its pathogenesis.

Wondfo<sup>®</sup> One Step HIV 1/2 Whole Blood/Serum/Plasma Test is a 3rd generation HIV immunoassay. The design of 3rd generation assays allows the detection of HIV specific IgG as well as IgM, which may occur early in infection.

#### PRINCIPLE OF THE PROCEDURE

Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test is a rapid immunochromatographic direct binding test for the visual detection of HIV antibodies in venous whole blood, fingerstick whole blood, serum or plasma samples in the diagnosis of HIV infection. Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test adopts double antigen sandwich method. When the specimen is added into the test device, the specimen is absorbed into the device by capillary action, mixes with the antigen-dye conjugate, and flows across the HIV antigen pre-coated membrane.

When the HIV antibody levels are at or above the detection limit of the test. HIV antibodies in the specimen bind to the antigen-dve conjugate and are captured by antigen immobilized in the test region (T) of the device. This produces a colored test band and indicates a positive result.

If no HIV antibodies are present or below the detection limit of the assay, no colored band will be visible in the test region (T) of the device. This indicates a negative result.

To serve as a procedure control, a colored line will appear at the control region (C), if the test has been performed properly.

#### WARNINGS AND PRECAUTIONS

1. This assay has been evaluated on EDTA, heparin, and sodium citrate. Other anticoagulants are excluded.

2.Read the Instructions for Use before using this product. The

instructions must be followed carefully as not doing so may result in inaccurate results.

- 3. Wear protective clothing such as laboratory coat, safety glasses and disposable gloves when handling specimens.
- 4.Wash hands thoroughly after use.Not intended for use in screening blood and tissue donors. This assay has not been evaluated for neonate or cord blood specimens.
- 5. This kit is for *in vitro* diagnosis of the infection of HIV only, this assay will not provide a diagnosis for any other disease.
- 6.Do not smoke, eat, drink, apply cosmetics or handle contact lenses in areas in which specimens are handled.
- 7. All specimens should be treated as potentially infectious.
- 8. EBV IgM positive specimens may cause erroneous results.
- 9. Discard test cassette, dropper, desiccant pouch after first useThe test can not be used more than once.
- 10. Cap and tightly the opened buffer after using.
- 11. Do not use the kit beyond the expiration date.
- 12. Do not use the kit if the pouch is punctured or not well sealed. Do not mix buffer /test devices from different kit lots.
- 13. Keep out of the reach of children.
- 14. All specimens and used-devices have infectious risks. The disposal process must follow the local infectious disposal law or laboratory rules.

#### CONTENT OF THE KIT

#### A. Kit components

Each individual sealed pouch contains one test cassette, one dropper and one desiccant pouch (for storage purposes only). Each test cassette contains one plastic cassette and one test strip.

The different article Number are as follows:

# Catalog No. W006-C4P2(2

Rev. A7

W006-P0045 

W006-P0046

W006-P0047

W006-P0048

W006-C4P2-F

W006-P0049

W006-P0050 \_\_\_\_\_

W006-P0051 

W006-P0052

# polyclonal antibody).

2.Sterile lancets

3.Alcohol Swabs

Components	Pouch Buffer Instruction	Alcohol swab Steri	Sterile lancet	Safety lancet (pcs)				
	( <b>pcs</b> )	(bottle)	( <b>pcs</b> )	(pcs)	( <b>pcs</b> )	18G	21G	23G
(25 Tests/Kit)	25	1	1	-	-	-	-	-
5 (25 Tests/Kit)	25	1	1	25	25	-	-	-
6 (25 Tests/Kit)	25	1	1	25	-	25	-	-
7 (25 Tests/Kit)	25	1	1	25	-	-	25	-
8 (25 Tests/Kit)	25	1	1	25	-	-	-	25
-F(40 Tests/Kit)	40	2	1	-	-	-	-	-
9 (40 Tests/Kit)	40	2	1	40	40	-	-	-
0 (40 Tests/Kit)	40	2	1	40	-	40	-	-
1 (40 Tests/Kit)	40	2	1	40	-	-	40	-
2 (40 Tests/Kit)	40	2	1	40	-	-	-	40

#### B. Reactive ingredients of main components

One test strip includes: Gold conjugate (HIV gp41/gp36 fusion recombinant antigen-gold colloid and rabbit IgG polyclonal 7. Specimen and test waste container antibody-gold colloid), Test line (HIV gp41 recombinant antigen and

#### MATERIALS REQUIRED BUT NOT PROVIDED

# For W006-C4P2(25 Tests/Kit) and W006-C4P2-F(40 Tests/Kit):

- 1. Specimen collection containers
- 4.Centrifuge (for serum/plasma specimen only)

## 5.Timer

Protective gloves

# HIV gp36 recombinant antigen) and Control line (Goat anti rabbit IgG For W006-P0046、W006-P0047、W006-P0048、W006-P0050、 W006-P0051 and W006-P0052:

- 1. Specimen collection containers
- 2. Centrifuge (for serum/plasma specimen only)
- 3 Timer
- 4. Protective aloves
- 5. Specimen and test waste container

### STORAGE AND OPERATING CONDITIONS

- 1. The kit and the unused buffer must be stored at 2°C-30°C before expiration date; the opened buffer should store at 2°C-30°C, no more than 8 weeks.
- 2. Use the kit within 1 hour once the pouch is opened.
- 3. Use the kit under the condition of the humidity from 20% to 90% and the temperature from  $10^{\circ}$  to  $30^{\circ}$ .
- 4. Recap and store the Buffer vial in the original container after use.
- 5. Shelf-life of the test devices and the Buffer are 24 months from the manufacturing date.
- 6. Keep away from sunlight, moisture, and heat.
- 7. Do not freeze.

#### COLLECTING AND PREPARING SPECIMENS

#### Whole blood – Collected by venipuncture

- 1. Using standard phlebotomy procedure, collect a venipuncture whole blood specimen using a blood collection tube with suitable anticoagulant (containing EDTA, citrate, or heparin).
- 2. It is recommended that specimens should be tested immediately. Do not leave the specimens at room temperature (10  $\degree$  to 30  $\degree$ ) within 4 hours. If the specimens are not tested immediately, they may be stored at 2°C-8°C for up to 7 days, only. It's not suitable to test the whole blood samples which have been stored at 2°C-8°C for more than 7 days.

#### Whole blood - Collected by fingerstick

- 1. Clean the area to be lanced with an alcohol pad. Allow the finger to dry thoroughly.
- 2. Using a lancet, puncture the inside of the finger pad. Apply gentle pressure beside the point of the puncture. Wipe away the first drop of blood with a sterile swab. Allow a new drop of blood to form. If blood flow is inadequate, the subject's finger may have to be gently massaged at the finger base to produce a droplet of sufficient volume.
- 3. Draw 10 microliter (µL) of finger blood with a capillary tube.
- 4. Whole blood specimens collected by fingerstick should be tested immediately.

#### Serum and Plasma

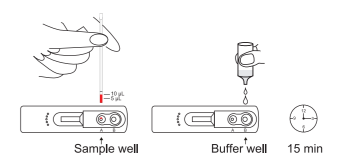
Rev. A7

- 1. Using standard phlebotomy procedure, collect a venipuncture whole blood specimen using a blood collection tube. If you need to collect plasma, please use a blood collection tube which contains suitable anticoagulant (EDTA, heparin, or sodium citrate).
- 2. Centrifuge whole blood and separate the plasma from red blood cell as soon as possible to avoid hemolysis.
- 3. Test should be performed immediately after the specimens have been collected. Do not leave the specimens at room temperature within 4 hours. Specimens should be stored at 2°C - 8°C if the test is to be run within 7 days of collection. If testing is delayed more than 7 days, the specimen should be frozen (-20 $^{\circ}$ C or colder). Bring specimens to room temperature (10  $\degree$  to 30  $\degree$ ) before testing. Frozen specimens must be completely thawed and mixed well prior to testing. Specimens should not be frozen and thawed repeatedly(Not more than 5 times). Only clear, non-hemolyzed specimens can be used.

#### TEST PROCEDURE

Equilibrate all specimens and the devices to room temperature (10)  $\sim$  30 °C ) before testing.

- 1. Remove a test cassette from the foil pouch by tearing the notch and place it on a level surface.
- 2. Use the specimen of either serum, plasma, or whole blood (collected by venipuncture): Slowly add 10µL (the second mark line of the dropper) of specimen to the sample well A, and then add 2 drops of buffer to the buffer well B.
- 3. Use the specimen of whole blood (collected by fingerstick): collecting the specimen by capillary tube (not provided), slowly add 10µL of specimen to the sample well A, and then add 2 drops of buffer to the buffer well B.
- 4. Incubate the cassette at room temperature (10  $\degree$   $\sim$  30  $\degree$ ), and read the result after 15 minutes, but not more than 30 minutes. Reading the test before 15 minutes or after 30 minutes may cause false result.



#### INTERPRETATION OF TEST RESULTS

#### Positive Result

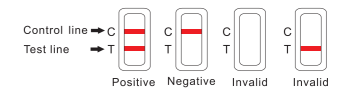
Rose-pink bands are visible in both the control region and the test region. A positive result indicates that the concentration of HIV1/2 antibodies in the sample is equal to or higher than the detection limit of the test.

#### Negative Result

A rose-pink band is visible in the control region. No color band appears in the test region. It indicates that the concentration of the HIV1/2 antibodies in the sample is below the detection limit of the test

#### Invalid Result

No visible band at all, or there is a visible band only in the test region but not in the control region, Repeat with a new test kit. If test still fails, please contact the distributor or the store, where you bought the product, with the lot number.



#### LIMITATIONS OF THE PROCEDURE

- 1. The kit is designed to detect antibodies against HIV-1 and HIV-2 in human serum, plasma, and whole blood,
- 2. The kit is a qualitative assay. It is not designed to determine the quantitative concentration of HIV antibodies.
- 3. The intensity of the T line does not necessarily correlate to the titer of antibody in the specimen.
- 4. The presence of the control line means only that liquid has flowed correctly. The control line will appear irrespective of whether a specimen is reactive or non-reactive.
- 5. As it is with any diagnostic procedure, a confirmed diagnosis should only be made after all clinical and laboratory findings have been evaluated.
- 6. A negative result with One Step HIV1/2 Whole Blood/Serum/ Plasma Test does not exclude the possibility of infection with HIV. A false negative result may occur in the following circumstances:
- Recent infection. Antibody response to a recent exposure may take several months to reach detectable levels.
- The test procedure has not been correctly followed.
- Antibodies to a variant strain of HIV1/2 in the patient do not react with specific antigens utilized in the assay configuration.
- Improper specimen handling.
- Failure to add sample.
- 7. A person who has antibodies to HIV-1 or HIV-2 is presumed to be infected with the virus, except that a person who has participated in an HIV vaccine study may develop antibodies to the vaccine and may or may not be infected with HIV. Clinical correlation is indicated with appropriate counseling, medical evaluation and possible additional testing to decide whether a diagnosis of HIV infection is accurate.

#### ROUBLESHOOTING

1. Buffer of less than 2 drops or more than 3 drops may cause incorrect results.

- 2. Specimen less than 5µL (the first mark line of the dropper) or more **2. Analytical performance study Cross reactivity** than 15µL (the third mark line of the dropper) may cause incorrect results.
- 3. The test result will be incorrect when adding the specimen and buffer in wrong position or sequence.

#### PERFORMANCE CHARACTERISTICS

1. Analytical performance study - Interfering substances

Summary of test results for determination of analytical specificity: potentially interfering substances.

#### A. Specimens of HIV-1 positive

Potentially interfering		Number of erroneous result		
substance	Number	Not spiked with anti-HIV positive specimen	Spiked with anti-HIV positive specimen	
Alcohol	6	0	0	
Haemoglobin	4	0	0	
Direct bilirubin	7	0	0	
Total bilirubin	17	0	0	
Triglyceride	17	0	0	
High-cholesterol	9	0	0	
Low density lipoprotein	15	0	0	
Rheumatic factor	25	0	0	
IgM gammopathies	10	0	0	
IgG gammopathies	2	0	0	
Pregnant women	31	0	0	
Systemic lupus erythematosus (SLE)	8	0	0	
Anti-nuclear antibodies (ANA)	8	0	0	
Anti-escherichia coli	2	0	0	
Total	161	0	0	

Summary of test results for determination of analytical speci- ficity: potentially cross-reacting unrelated infections and diseases.

Potentially cross-reacting substance	Number	Number of erroneous resul
Malaria	27	0
Epstein-Barr virus immunoglobulin(EBV lgM) <sup>1</sup>	5	2
Epstein-Barr virus immunoglobulin(EBV IgM) <sup>2</sup>	60	0
Influenza antibody	13	0
Cytomegalovirus immunoglobulin M(CMV IgM)	5	0
Syphilis	5	0
Herpes simples virus(HSV)	5	0
Anti-HBc	15	0
Ant-HBs	15	0
Anti-HCV	15	0
Anti-HTLV 1/2	15	0
Anti-HEV	10	0
Total	185	2

1. Testing in the third party Institute of Tropical Medicine

2. Testing in Guangzhou Wondfo Biotech Co.,Ltd.

#### 3. Analytical performance study - Analytical sensitivity

A total of 45 HIV seroconversion panels were tested, 25 of which were tested by Wondfo and 20 were tested by a third party institution. Among the 25 HIV seroconversion panels tested by Wondfo with a commercially available WHO pregualified HIV ELISA reagent as reference assay, HIV antibody in 6 panels was detected by Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test earlier than that by the ELISA; HIV antibody in 2 panels that detected by Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test were later than that by the ELISA; and HIV antibody in 17 panels were detected by both assays at the same bleeding.

For the 20 HIV seroconversion panels tested by the third party institution, 4 CE marked HIV ELISA reagents were set as reference assay. From the 111 seroconversion panel members, the One Step HIV 1/2 Whole Blood/Serum/Plasma Test detected 18 samples more than the least sensitive CE marked antibody test and 4 samples less than the most sensitive CE marked antibody test.

## 4. Analytical performance study - HIV-1 subtypes positive specimens

The HIV-1 Worldwide Performance Panel, the WHO International Standard HIV (antibody), 1st International Panel (NIBSC code: 02/210) and 50 specimens with various subtypes (10 specimens for each of subtypes A1, B, C, CRF02 AG and G) were tested by Wondfo. And 40 specimens with various subtypes were tested by a third party institution, including 3 specimens for each of following subtypes: C, CRF01 AE, CRF02 AG, CRF06 cpx, CRF36 cpx, D, G, group O, H, J, and K, 2 specimens for subtypes A, F1 and F2, and 1 specimen for subtype A1.

Wondfo One Step HIV1/2 Whole Blood/Serum/Plasma Test can detect HIV-2 antibodies and the following subtypes of HIV-1 antibodies: Group O, subtype A, subtype A1, subtype B, subtype C, subtype D, subtype F1, subtype F2, subtype H, subtype J, subtype K, subtype G, CRF01-AE, CRF02\_AG, CRF06\_cpx, CRF36\_cpx and CRF02-AG.

#### 5. Clinical performance study - Diagnostic sensitivity

Summary of results of a clinical study to determine diagnostic sensitivity.

Specimens type	No. of specimens	False negative	Sensitivity	95% Confidence Interval
Serum	456	0	100%	(99.18, 100)
Plasma	620	0	100%	(99.38, 100)
Whole venous blood	100	0	100%	(96.30, 100)
Total	1176	0	100%	(99.67, 100)

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Specimens type ------Plasma

specificity.

Specimens type	No. of specimens	False positive	Specificity	95% Confidence Interval
Serum	331	0	100.00%	(98.85, 100)
Plasma	1904	1	99.95%	(99.70, 99.99)
Whole venous blood	500	0	100.00%	(99.24, 100)
Total	2735	1	99.96%	(99.79, 99.99)

#### LIST OF REFERENCES

- 3. Gale R. Burstein, Jonathan Pincus et al. A rapid review of rapid HIV antibody tests. Current Infectious Disease Reports, Volume 8, Number 2, March 2006, Pages 125-131.

- 5. Holm-Hansen C, Constantine NT, Haukenes G. Detection of antibodies to HIV in homologous sets of plasma, urine and oral mucosal transudate sample susing rapid assays in Tanzania[J]. ClinDiagnVirol, 1993,1(4): 207-214.

#### B. Specimens of HIV-2 positive

## SYMBOLS KEY

Caution

No. of specimens	False negative	Sensitivity	95% Confidence Interval
100	0	100%	(96.30, 100)

#### 6. Clinical performance study - Diagnostic specificity

Summary of results of a clinical study to determine diagnostic

- 1. Janssen R. S. etal. New testing strategy to detect early HIV-1 infection for use in incidence estimates and for clinical and prevention purposes. JAMA(1998) 280(1): 42-48.
- 2. CDC. Update: HIV Counseling and Testing using Rapid Tests please contact us by phone, e-mail or in writing. United States, 1995. MMWR 1998; 47(11).
- 4. Bernard M. Branson. State of the Art for Diagnosis of HIV Infection. Clinical Infectious Diseases 2007; 45: S221-S225.

In vitro diagnostic medical device	Consult Instructions for Use	Expiry Date
Content sufficient for < n > tests	Date of manufacture	Keep dry
LOT Batch code	2°C Temperature limit	Keep away from sunlight
Indicates the device manufacturer	Do not re-use	REF Product code/ Catalogue number

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Any complaints, questions, problems, suggestions or comments,

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