

**WHO Prequalification of In Vitro Diagnostics Programme
PUBLIC ASSESSMENT REPORT**

**Product: Bioline HIV 1/2 3.0
Number: PQDx 0027-012-00**

Bioline HIV 1/2 3.0 with product codes 03FK10, 03FK16, 03FK17, and 03FK22, manufactured by Abbott Diagnostics Korea Inc., Rest-of-World regulatory version was accepted for the WHO list of prequalified in vitro diagnostics and was listed on 20 May 2013.

Summary of Prequalification Assessment for the Bioline HIV 1/2 3.0

	Date	Outcome
Status on PQ list	20 May 2013	listed
Dossier assessment	11 August 2011	MR
Product performance evaluation	05 April 2013	MR

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 summarised in the following table, and details of each amendment are provided below.

Version	Summary of amendments and change request reference where applicable.	Date of report amendment
2.0	Review of text updates in the public report	2013
3.0	Change of inner structure of device to capture a reasonable excess volume of buffer that might be added by the end user, thereby reducing the risk of overflow of buffer.	2013

4.0	Addition of a product code, which corresponds to the IVD supplied with the new safety lancets. Product code is 03FK17, 25 tests per kit, with 25 safety lancets	23 February 2017
5.0	Change of product name from SD BIOLINE HIV-1/2 3.0 to Bioline HIV 1/2 3.0 and change of manufacturer's name from Standard Diagnostics, Inc to Abbott Diagnostics Korea Inc.	20 August 2020
6.0	Accessory labelling changes according to accessory manufacturer information change (Manufacturer name change and EU representative address change) (PQC-IVD-2024-0033). Addition of new product specification of Bioline HIV 1/2 3.0 for Ghana. This change introduces a new packaging specification for the Ghanaian market, designated 03FK22, which includes 5 buffers per kit and is tailored for the National AIDS Control Program (NACP) of the Ministry of Health, Ghana. In line with this new specification, updates have also been made to the IFU and packaging (PQC-IVD-2025-0068).	20 March 2026
7.0	Addition of a capillary tube label (PQC-IVD-2025-0118).	15 May 2026

Intended use¹:

According to the claim of intended use from Abbott Diagnostics Korea Inc, "*Bioline HIV 1/2 3.0 kit is a rapid, qualitative test for the detection of antibodies to all isotypes (IgG, IgM, IgA) specific to HIV-1 and HIV-2 simultaneously in human serum, plasma or whole blood. The Bioline HIV 1/2 3.0 kit is intended only for professional use and for in vitro diagnostic use. This test may not be suitable for diagnosis of early infection or blood donation screening. Positive samples should be confirmed by a supplemental assay such as ELISA or Western Blot test*".

¹ This product is one that uses Protein A to detect human IgG antibodies. Protein is also able to detect other classes of human antibody (IgA, IgD, IgE and IgM) but not as reliably as it does IgG. This product has been prequalified with respect to its ability to detect human IgG antibodies. Any claim to detect other types of antibodies on this kind of product has not been validated based on WHO prequalification requirements.

Test kit contents:

	30T/kit (product code 03FK10)	25T/kit (product code 03FK16)	25T/kit (product code 03FK17)	25T/kit (product code 03FK22)
Test cassettes individually packed in foil pouch with a desiccant	30 test devices	25 test devices	25 test devices	25 test devices
Assay diluent dispensed in plastic bottle	1 x 4 mL/bottle	1 x 4 mL/bottle	1 x 4 mL/bottle	5 x 4 mL/bottle
Specimen transfer devices Capillary pipettes (20µl)	N/A	25 units of 20 µL	25 units of 20 µL	25 units of 20 µL
Lancets Disposable, sterilized	N/A	25 units	25 units (safety lancet)	25 units (sterile lancet)
Alcohol swabs Disposable	N/A	25 units	25 units	25 units
Instructions for use	1 copy	1 copy	1 copy	5 copies

Storage:

The test kit must be stored between 1 and 30 °C.

Shelf-life upon manufacture²:

24 months.

Dossier assessment

Abbott Diagnostics Korea Inc submitted a product dossier for Bioline HIV 1/2 3.0 as per the "Instructions for compilation of a product dossier" (PQDx_018 v1). The information submitted in the product dossier was reviewed by WHO staff and external experts (assessors) appointed by WHO.

Based on the assessment of the product dossier, Bioline HIV 1/2 3.0 meets WHO prequalification requirements.

² The assigned device shelf-life is based on stability data generated from the date of manufacture. The finished goods shelf-life, calculated from the date of packaging completion, may be shorter depending on the time elapsed between manufacture and final packaging of the device.

Manufacturing site inspection

The inspection of the manufacturing site(s) was conducted to assess whether the manufacturer's quality management system (QMS) and manufacturing practices are in alignment with:

- (i) applicable international standards, such as ISO 13485 (Medical devices – Quality management systems – Requirements for regulatory purposes);
- (ii) the manufacturer's own documented procedures and quality requirements; and
- (iii) other relevant international standards and guidelines applicable to in vitro diagnostic (IVD) medical devices. The WHO's Public Inspection Reports are accessible at:

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

Product performance evaluation

Bioline HIV 1/2 3.0 was evaluated by WHO at the Institute of Tropical Medicine, Antwerp, Belgium, in the last quarter of 2012 using serum/plasma specimens. From this evaluation, we drew the following conclusions:

Bioline HIV 1/2 3.0 is a lateral flow immunochromatographic assay for the discriminatory detection HIV-1 and HIV-2 antibodies in human serum/plasma and whole blood. A volume of 10 µL of serum/plasma or 20 µL of whole blood is required 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 is performed visually i.e. subjectively read.

In this limited evaluation on a panel of 1118 clinically-derived specimens, we found an initial sensitivity (95% CI) of 99.8% (98.8% - 100%) and an initial specificity (95% CI) of 99.9% (99.2% - 100%) compared to the reference assays (Vironostika HIV Ag/ab [bioMérieux] and Enzygnost Anti-HIV 1/2 in parallel; followed by INNO-LIA HIV H/II Score [Innogenetics]). The final sensitivity (95% CI) was 100% (99.2% - 100%) and the final specificity (95% CI) was 99.9% (99.2% - 100%) compared to the reference assays. Lot to lot variation observed was within the acceptance criteria.

Bioline HIV 1/2 3.0 was unable to discriminate between HIV-1 and HIV-2 for seven HIV-2 specimens, and 22 HIV-1 specimens (6.3% of 460 HIV positive specimens), as two test bands of equal intensity were observed.

For eight seroconversion panels, Bioline HIV 1/2 3.0 detected on average 0.125 specimens later than the benchmark assay (Enzygnost Anti-HIV 1/2 Plus [Siemens Healthcare Diagnostics]).

For the mixed titer panel, Bioline HIV 1/2 3.0 correctly classified all specimens. For the 1st International Reference Panel for anti-HIV [NIBSC code 02/210], Bioline HIV 1/2 3.0 detected all HIV-1 subtypes tested (HIV-1 A, HIV-1 B, HIV-C, HIV-1 CRF01_AE, and HIV-2).

In this study, 0% of the results were recorded as indeterminate. Results were interpreted independently by three technicians; the overall inter-reader variability was 1.9% (0.2% for HIV-1 band, 1.8% for HIV-2 band). The invalid rate was 0%.

Based on these results, the performance evaluation for Bioline HIV 1/2 3.0 meets the WHO prequalification requirements.

Labelling review

The labelling submitted for Bioline HIV 1/2 3.0 was reviewed by WHO staff and external technical experts appointed by WHO. The review evaluated the labelling for clarity and consistency with the information submitted in the product dossier, alignment with international guidance and standards, and suitability for the intended users and settings in WHO Member States, including low- and middle-income countries.

The table below provides traceability of the labelling documents reviewed during the assessment, including document titles, version numbers, approval dates, and control identifiers.

Controlled Labelling References

03FK10

Document Type	Document Title	Version / Revision	Date Approved	Controlled Document No.
Outer box artwork	Package	A0	2019-11-18	03FK10-40-A0
Pouch / Device label	Device pouch label	A0	2019-11-01	FP26-A0
Assay diluent labels	Assay diluent labels	A0	2019-10-25	FB07-A0
Instructions for Use (IFU)	IFU	A0	2020-03-04	03FK10/03FK16-04-A0

03FK16

Document Type	Document Title	Version / Revision	Date Approved	Controlled Document No.
Outer box artwork	Package	A0	2019-11-18	03FK16-40-A0
Pouch / Device label	Device pouch label	A0	2019-11-01	FP26-A0
Assay diluent labels	Assay diluent labels	A0	2019-10-25	FB07-A0
Capillary Tube	40005438	2	2024-05-21	D10151501
Capillary Tube	40001927	0	2026-03-23	D10167079
Sterile non-safety lancet	P65-008	0	2023-11-10	D10151501
Sterile non-safety lancet	40006487	0	2024-10-01	D10151501
IFU	IFU	A0	2020-03-04	03FK10/03FK16-04-A0

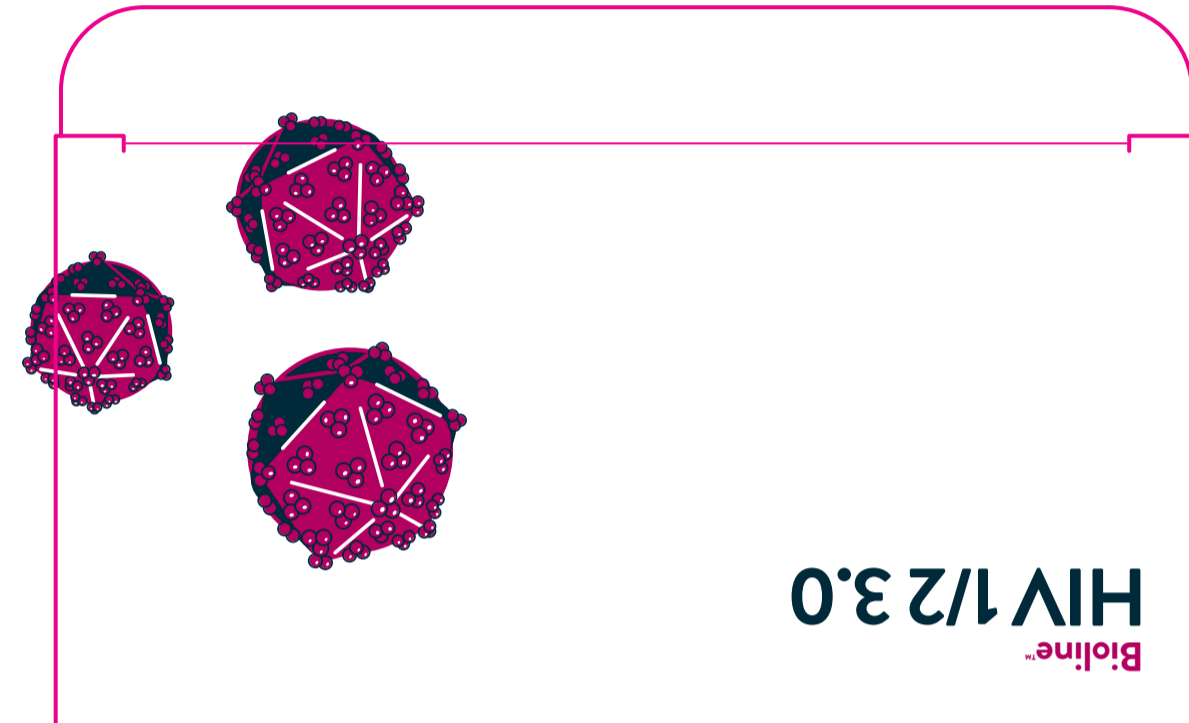
03FK17

Document Type	Document Title	Version / Revision	Date Approved	Controlled Document No.
Outer box artwork	Package	A0	2019-11-20	03FK17-40-A0
Pouch / Device label	Device pouch label	A0	2019-11-01	FP26-A0
Assay diluent labels	Assay diluent labels	A0	2019-10-25	FB07-A0
Capillary Tube	40005438	2	2024-05-21	D10151501
Capillary Tube	40001927	0	2026-03-23	D10167079
Sterile safety lancet	P65-013/000/001	0	2023-11-10	D10151501
Sterile safety lancet	P65-013/003	0	2023-11-10	D10151501
Sterile safety lancet	P65-013/006	0	2023-11-10	D10151501
IFU	IFU	A0	2021-09-09	03FK17-04-A0

03FK22



Document Type	Document Title	Version / Revision	Date Approved	Controlled Document No.
Outer box artwork	Package	A0	2023-03-23	03FK22-40-GH-A0
Pouch / Device label	Device pouch label	A0	2019-11-01	FP26-A0
Assay diluent labels	Assay diluent labels	A0	2019-10-25	FB07-A0
Capillary Tube	40005438	2	2024-05-21	D10151501
Capillary Tube	40001927	0	2026-03-23	D10167079
Sterile safety lancet	P65-008	0	2023-11-10	D10151501
Sterile safety lancet	40006487	0	2024-10-01	D10151501
IFU	IFU	A0	2023-03-23	03FK22-01-EN-A0

Labels

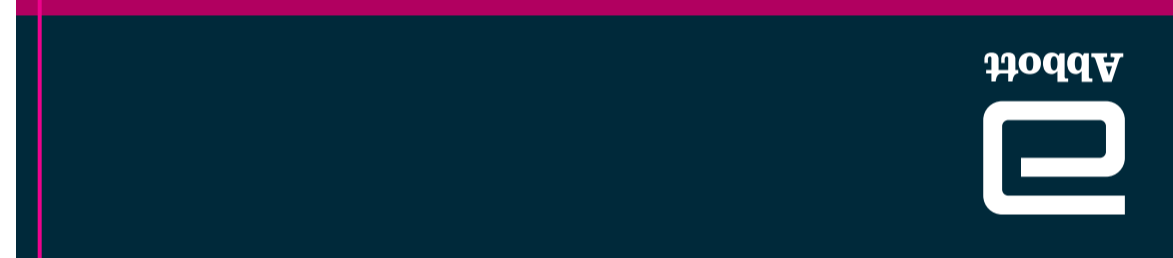


Bioline[®]
HIV 1/2 3.0

PRODUCT - Bioline™ HIV 1/2 3.0 30T
 SIZE - 152 X 115 X 70 MM

 PMS 303 C
 PMS 227 C

PN. - 03FK10-40-A0
 REV. DATE - 2019.10.11
 CREATOR - THINKBOX



EN Rapid, immunochromatographic test for the detection of anti-HIV 1/2 in human serum, plasma or whole blood

FR Test rapide par immunochromatographie pour la détection de anti-HIV 1/2 en sérum humain, plasma ou sang total

ES Rápido, test inmunocromatográfico para la detección de anti-HIV 1/2 en suero humano, plasma o sangre entera

PT Teste rápido, imunocromatográfico, para a deteção do anti-HIV 1/2 em amostra de soro, plasma ou sangue total humano

CONTENTS:

- 30 Test devices with desiccant in individual foil pouches
- Assay diluent (1 x 4 ml/vial)
- 1 Instructions for use

CONTENU :

- 30 dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium
- Diluant du test (1 x 4 ml/flacon)
- 1 mode d'emploi

CONTENIDO:

- 30 dispositivos de prueba con desecante en bolsas de papel de aluminio individuales
- Diluyente del ensayo (1 x 4 ml/vial)
- 1 instrucciones de uso

CONTEÚDO:

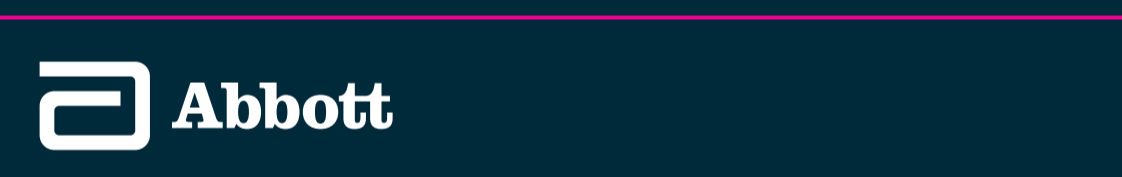
- 30 dispositivos de teste com desecante em bolsas de alumínio individuais
- Diluente de ensaio (1 x 4 ml/frasco)
- 1 Instruções de utilização

03FK10-40-A0



Bioline[®]
HIV 1/2 3.0

PRINT AREA



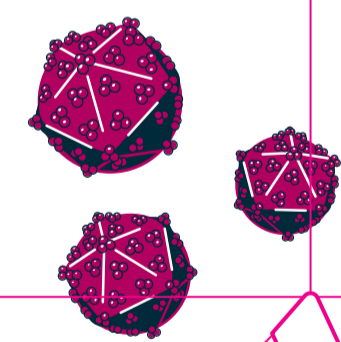
Bioline[®]
HIV 1/2 3.0







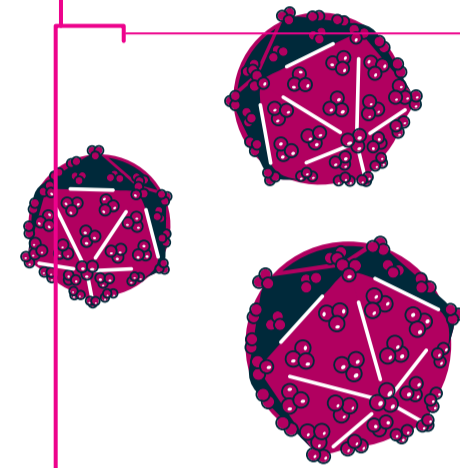


Bioline[®]
HIV 1/2 3.0

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Bioline™
HIV 1/2 3.0



Bioline™
HIV 1/2 3.0



EN
Rapid, immunochromatographic test for the detection of anti-HIV 1/2 in human serum, plasma or whole blood

FR
Test rapide par immunochromatographie pour la détection de anti-HIV 1/2 dans le sérum humain, plasma ou sang total

ES
Rápido, test inmunocromatográfico para la detección de anti-HIV 1/2 en suero humano, plasma o sangre entera

PT
Teste rápido, imunocromatográfico, para a deteção do anti-HIV 1/2 em amostra de soro, plasma ou sangue total humano

CONTENTS:
• 25 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL/vial)
• 25 Capillary pipettes (20 µL)
• 25 Sterile lancets, 25 Alcohol swabs
• 1 Instructions for use

CONTENU :
• 25 dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium
• Diluant du test (1 x 4 mL/flacon)
• 25 pipettes capillaires (20 µL)
• 25 lancettes stériles, 25 compresses d'alcool
• 1 mode d'emploi

CONTENIDO:
• 25 dispositivos de prueba con desecante en bolsas de papel de aluminio individuales
• Diluyente del ensayo (1 x 4 mL/vial)
• 25 pipetas capilares (20 µL)
• 25 lancetas estériles, 25 hisopos con alcohol
• 1 instrucciones de uso

CONTEÚDO:
• 25 dispositivos de teste com dessecante em bolsas de papel de alumínio individuais
• Diluente de ensaio (1 x 4 mL/frasco)
• 25 pipetas capilares (20 µL)
• 25 lancetas estériles, 25 aragotes com álcool
• 1 Instrukções de utilização

03FK16-40-A0

Bioline™
HIV 1/2 3.0

PRINT AREA

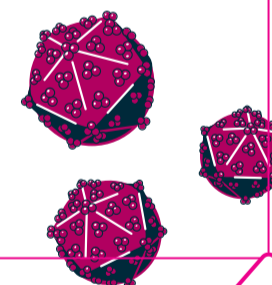
Bioline™
HIV 1/2 3.0



1°C - 30°C
34°F - 86°F



03FK16



Bioline™
HIV 1/2 3.0

Abbott Diagnostics Korea Inc.
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Gyeonggi-do, 17099, Republic of Korea
abbott.com/kr

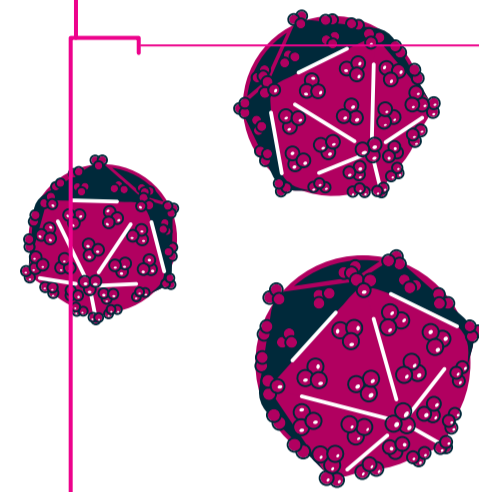
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PRODUCT - Bioline™ HIV 1/2 3.0, 25T

SIZE - 152 X 115 X 58 MM

PMS 303 C
PMS 227 C

PN. - 03FK16-40-A0
REV. DATE - 2019.10.11
CREATOR - THINKBOX



Bioline[®] HIV 1/2 3.0



Bioline[®]
HIV 1/2 3.0

EN
Rapid, immunochromatographic test
for the detection of anti-HIV 1/2 in
human serum, plasma or whole blood

FR
Test rapide par immunochromatographique
pour la détection de anti-HIV 1/2 dans le
sérum humain, plasma ou sang total

ES
Test rápido, inmunocromatográfico para
la detección de anti-HIV 1/2 en suero
humano, plasma o sangre entera

PT
Teste rápido, imunocromatográfico, para
a deteção de anti-HIV 1/2 em amostra
de soro, plasma ou sangue total humano

CONTENTS:
• 25 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 ml/vial)
• 25 Capillary pipettes (20 µl),
25 Safety lancets,
25 Alcohol swabs
• 1 Instructions for use

CONTENU :
• 25 dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium
• Diluant du test (1 x 4 ml/flacon)
• 25 pipettes capillaires (20 µl),
25 lancettes de sécurité,
25 compresses d'alcool
• 1 mode d'emploi

CONTENIDO:
• 25 dispositivos de prueba con desecante en bolsas de papel de aluminio individuales
• Diluyente del ensayo (1 x 4 ml/vial)
• 25 pipetas capilares (20 µl),
25 lancetas de seguridad,
25 hisopos con alcohol
• 1 instrucciones de uso

CONTEÚDO:
• 25 dispositivos de teste com dessecante em bolsas de alumínio individuais
• Diluente de ensaio (1 x 4 ml/frasco)
• 25 pipetas capilares (20 µl),
25 lancetas de segurança,
25 saragatas com álcool
• 1 Instruções de utilização

03FK17-40-A0

PRODUCT - Bioline™ HCV 25T,
SAFETY LANCET

SIZE - 152 X 115 X 75 MM

PMS 303 C
PMS 227 C

CAT NO. - 03FK17-40-A0
REV. DATE - 2019. 10. 11
CREATOR - THINKBOX



Bioline[®]
HIV 1/2 3.0

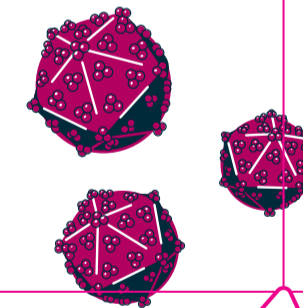
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Bioline[®]
HIV 1/2 3.0



1°C 30°C 34°F 86°F 25 1 2 IVD 100% REF 03FK17



Bioline[®]
HIV 1/2 3.0

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MINISTRY OF HEALTH, GHANA - NOT FOR SALE



Bioline™ HIV 1/2 3.0

Abbott

Abbott Bioline™
HIV 1/2 3.0

<p>EN Rapid, immunochromatographic test for the detection of anti-HIV 1/2 in human serum, plasma or whole blood</p> <p>CONTENTS:</p> <ul style="list-style-type: none"> • 25 Test devices with desiccant in individual foil pouches • Assay diluent (5 x 4 ml/vial) • 25 Capillary pipettes (20 µl) • 25 Sterile lancets, 25 Alcohol swabs • 5 Instructions for use <p>03FK22-40-GH-A0</p>	<p>FR Test rapide par immunochromatographique pour la détection de anti-HIV 1/2 dans le sérum humain, plasma ou sang total</p> <p>CONTENU :</p> <ul style="list-style-type: none"> • 25 dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium • Diluant du test (5 x 4 ml/flacon) • 25 pipettes capillaires (20 µl) • 25 lancettes stériles, 25 compresses d'alcool • 5 mode d'emploi 	<p>ES Rápido, test inmunocromatográfico para la detección de anti-HIV 1/2 en suero humano, plasma o sangre entera</p> <p>CONTENIDO:</p> <ul style="list-style-type: none"> • 25 dispositivos de prueba con desecante en bolsas de papel de aluminio individuales • Diluyente del ensayo (5 x 4 ml/vial) • 25 pipetas capilares (20 µl) • 25 lancetas estériles, 25 hisopos con alcohol • 5 instrucciones de uso 	<p>PT Teste rápido, imunocromatográfico, para a deteção do anti-HIV 1/2 em amostra de soro, plasma ou sangue total humano</p> <p>CONTEÚDO:</p> <ul style="list-style-type: none"> • 25 dispositivos de teste com dessecante em bolsas de alumínio individuais • Diluente de ensaio (5 x 4 ml/frasco) • 25 pipetas capilares (20 µl) • 25 lancetas estériles, 25 anagetas com álcool • 5 instruções de utilização
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PRODUCT - Bioline™ HIV 1/2 3.0, 25T , Ghana SIZE - 152 x 115 x 70mm	<div style="display: flex; justify-content: space-between;"> <div style="width: 40%;"> <div style="width: 100%; height: 10px; background-color: #003366; margin-bottom: 2px;"></div> PMS 303 C </div> <div style="width: 40%;"> <div style="width: 100%; height: 10px; background-color: #990033; margin-bottom: 2px;"></div> PMS 227 C </div> </div>	PN - 03FK22-40-GH Rev - A0 CREATOR - Jamie Lee REV.DATE - 2023.01.27
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<p style="font-size: 24px; margin: 0;">Bioline™ HIV 1/2 3.0</p> <div style="border: 1px solid #990033; padding: 10px; width: 80px; margin: 10px auto;"> <p style="font-size: 24px; margin: 0; color: #990033;">PRINT AREA</p> </div>	<p style="font-size: 24px; margin: 0;">Bioline™ HIV 1/2 3.0</p>  <div style="display: flex; justify-content: center; align-items: center; margin-top: 10px;"> <div style="font-size: 8px; margin-right: 5px;"> 1°C 30°C 34°F 86°F </div> <div style="font-size: 8px; margin-right: 5px;"> 25 </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; margin-right: 5px;"> </div> <div style="font-size: 8px; 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PRODUCT - Biline™

Assay diluent label

SIZE - 50 X 16 MM



PMS 303 C



PMS 2925 C

CAT NO. - FB07-A0

REV. DATE - 2019.10.21

CREATOR - SCOTT KIM

	ITEM HIV 1/2 3.0
Biline™ ASSAY DILUENT	VOL 4ml/vial
IVD 1°C → 30°C	LOT XXXXXXXXX
<small>Abbott Diagnostics Korea Inc. FB07-A0</small>	Y YY.MM.DD

40005438_Capillary Tube (20ul)25PCS_60×60



20µl Capillary Tube

Code No. : 40005438
LOT No. :
Quantity : 25PCS
MFG Date :

HLB Life Science Co.,Ltd
101-908, Digital Empire11, 88, Sinwon-ro, Yeongtong-gu
Suwon-si, Gyeonggi-do, 16681, Republic Korea
(01) 08809899620106



Cmc Medical Devices & Drugs S.L.
C/ Horacio Lengo n18 C.P 29006 Málaga-Spain



20µl Capillary Tube

Code No. : 40005438
LOT No. :
Quantity : 25PCS
MFG Date :

HLB Life Science Co.,Ltd
101-908, Digital Empire11, 88, Sinwon-ro, Yeongtong-gu
Suwon-si, Gyeonggi-do, 16681, Republic Korea
(01) 08809899620106



Cmc Medical Devices & Drugs S.L.
C/ Horacio Lengo n18 C.P 29006 Málaga-Spain



Capillary tube

Model: Single marked line

Specification: 20µl



FOR PROFESSIONAL USE ONLY



Jiangsu Changfeng Medical Industry Co.,ltd

Address: 189 Tongda Road, Touqiao Town, Yangzhou, China

Tel:+86-13816762056

E-mail: Jacky. cao@changfeng-medical.com

EC

REP

Llins Service & Consulting GmbH

Address: Obere Seegasse 34/2, 69124, Heidelberg, Germany

Email: info@llins-service.com



(01) XXXXXXXXXXXXXXXX

(11) YYMMDD

(17) XXXXXX

(10) XXXXXXXX



YYYY-MM-DD



YYYY-MM-DD



XXXXXXXXXX

Made in China

A/0



PANTONE Reflex Blue C
Shandong
Size:50X40

Sterile Lancet for single use

LOT LOT NO./Type : 2305267303 / 28G
EXP. DATE : 2028-06-04
QTY : 25pcs

STERILE R **CE** 0123 

010004051700443617288004193205207101

EC REP Shanghai International Holding Corp. GmbH (Europe)
Eiffelstrasse 60, 20537 Hamburg, Germany

Shandong Lianfa Medical Plastic Products Co., Ltd.
No. 3 Shuangshihai Sanjian Road, 250200 Zhangqiu City, Jinan, Shandong
PEOPLE'S REPUBLIC OF CHINA

PANTONE Reflex Blue C
Shandong
Size:50X40

Safety Lancet










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 EXP. DATE : 2028-05-09
 QTY : 25pcs

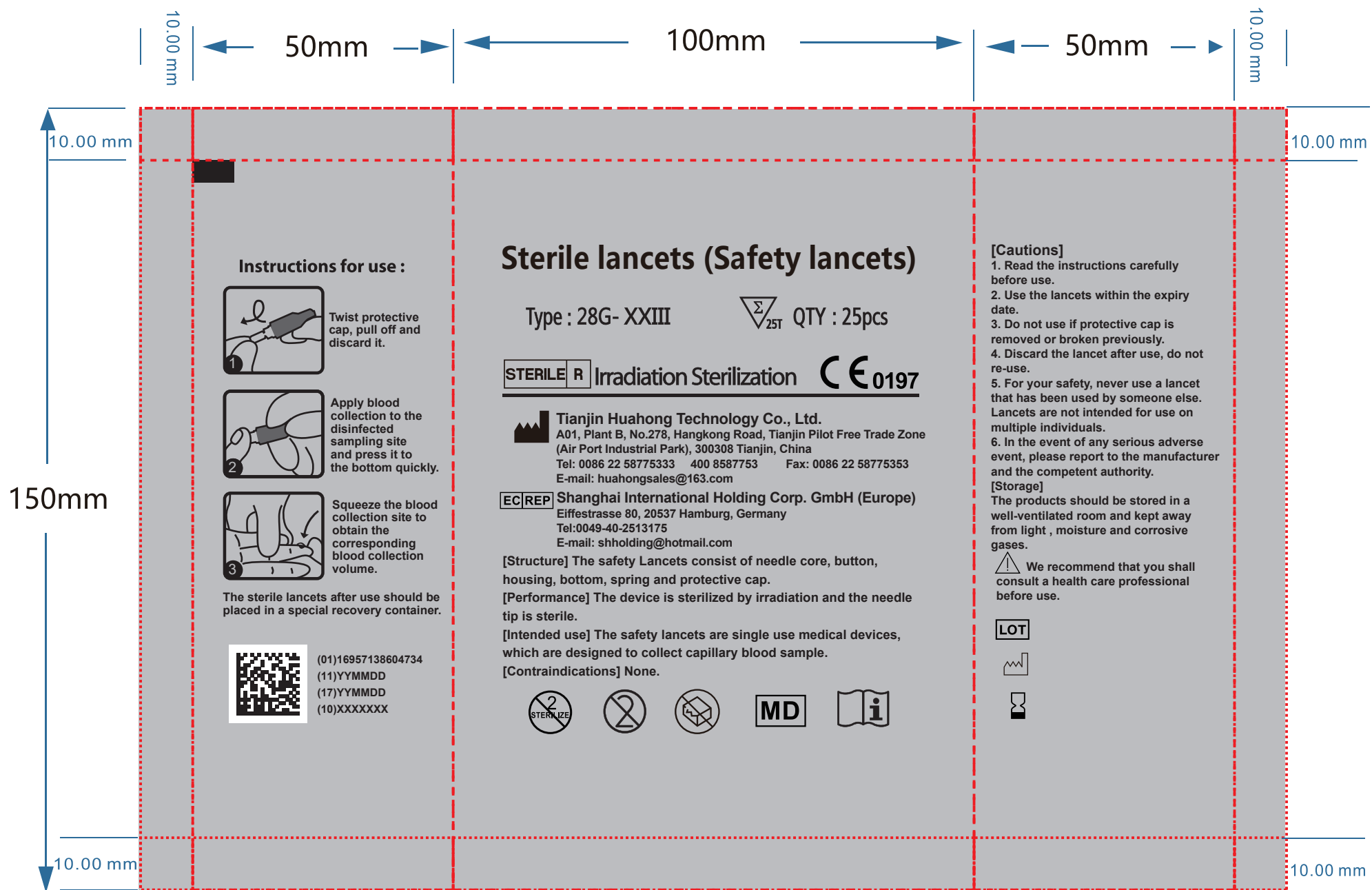


STERILE R  **0123** (010004851700104723000001023032692)

EC REP Shanghai International Holding Corp. GmbH (Europe)
Eiffelstrasse 60, 20537 Hamburg, Germany

 **Shandong Lianfa Medical Plastic Products Co., Ltd.**
No. 3 Shuangshihai Sanjian Road 250200 Zhangou City, Jinan, Shandong
PEOPLE'S REPUBLIC OF CHINA

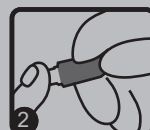
	10mm	33.75mm	67.5mm	33.75mm	10mm
10mm	<div style="display: flex; justify-content: space-between;"> <div style="width: 20%;">   <p>(01)16957138602815 (11)YYMMDD (17)YYMMDD (10)XXXXXXXX</p> <p>LOT</p>   <p>EC REP Shanghai International Holding Corp. GmbH (Europe) Eiffestrasse 80, 20537 Hamburg, Germany Tel: 0049-40-2513175 E-mail: shholding@hotmail.com</p> </div> <div style="width: 50%; text-align: center;"> <p>Twist-off lancet (use with lancing device) 25 pcs (28G I)</p> <p>[Instructions]</p> <ol style="list-style-type: none"> Unscrew the lancing device cap from the body of the lancing device. Insert a sterile lancet into the lancing device firmly. Twist off the protective cap of the lancet. Carefully screw the lancing device cap back to the lancing device. Adjust the puncture depth. Put the lancing device against the side of the finger and press the release button to prick your fingertip. Apply a drop of blood to test strip. Recap lancet and discard it in appropriate container. Place the lancing device cap back to the lancing device. <p>[Structure] The Twist-off Lancets is consist of needle, main body and protective cap.</p> <p>[Performance] The device is sterilized by irradiation.</p> <p>[Intended use] The twist-off lancet (use with lancing device) is a single use, sterile medical device, which is designed for use of micro blood sampling puncture to obtain capillary blood samples from the fingertip.</p> <p>[Contraindications] None.</p> <p>[Shelf life] 5 years.</p> <p>[Storage] The products should be stored in a well-ventilated room and kept away from light, moisture and corrosive gases.</p> <p>Tianjin Huahong Technology Co., Ltd. A01, Plant B, No.278, Hangkong Road, Tianjin Pilot Free Trade Zone (Air Port Industrial Park), 300308 Tianjin, China Tel: 0086 22 58775333 400 8587753 Fax: 0086 22 58775353 E-mail: huahongsales@163.com</p> </div> <div style="width: 25%;"> <p>[Cautions]</p> <ol style="list-style-type: none"> Read the instructions carefully before use. Use the lancets within the expiry date. Do not use if protective cap is removed or broken previously. Do not store lancet in lancing device, discard lancet after use, do not re-use. For your safety, please do not use a lancet that has been used by someone else. Lancets are not intended for use on multiple individuals. In the event of any serious adverse event, please report to the manufacturer and the competent authority. <p> The lancet and lancing device must be compatible with each other. When you use Huahong's sterile lancets with other manufacturers' lancing devices, please be sure that these products are compatible. We recommend that you shall consult a health care professional before use.</p> <p>STERILE R</p>    <p> 0197</p> </div> </div>				
75mm					
10mm					



Instructions for use :



Twist protective cap, pull off and discard it.



Apply blood collection to the disinfected sampling site and press it to the bottom quickly.



Squeeze the blood collection site to obtain the corresponding blood collection volume.

The sterile lancets after use should be placed in a special recovery container.



(01)16957138604734
 (11)YYMMDD
 (17)YYMMDD
 (10)XXXXXX

Sterile lancets (Safety lancets)

Type : 28G- XXIII

QTY : 25pcs

STERILE R Irradiation Sterilization CE 0197

Tianjin Huahong Technology Co., Ltd.
 A01, Plant B, No.278, Hangkong Road, Tianjin Pilot Free Trade Zone
 (Air Port Industrial Park), 300308 Tianjin, China
 Tel: 0086 22 58775333 400 8587753 Fax: 0086 22 58775353
 E-mail: huahongsales@163.com

EC/REP Shanghai International Holding Corp. GmbH (Europe)
 Eiffestrasse 80, 20537 Hamburg, Germany
 Tel:0049-40-2513175
 E-mail: shholding@hotmail.com

[Structure] The safety Lancets consist of needle core, button, housing, bottom, spring and protective cap.

[Performance] The device is sterilized by irradiation and the needle tip is sterile.

[Intended use] The safety lancets are single use medical devices, which are designed to collect capillary blood sample.

[Contraindications] None.



[Cautions]

1. Read the instructions carefully before use.
2. Use the lancets within the expiry date.
3. Do not use if protective cap is removed or broken previously.
4. Discard the lancet after use, do not re-use.
5. For your safety, never use a lancet that has been used by someone else. Lancets are not intended for use on multiple individuals.
6. In the event of any serious adverse event, please report to the manufacturer and the competent authority.

[Storage]

The products should be stored in a well-ventilated room and kept away from light , moisture and corrosive gases.

We recommend that you shall consult a health care professional before use.

LOT



10.00 mm

10.00 mm

150mm

10.00 mm

10.00 mm

10.00 mm

10.00 mm

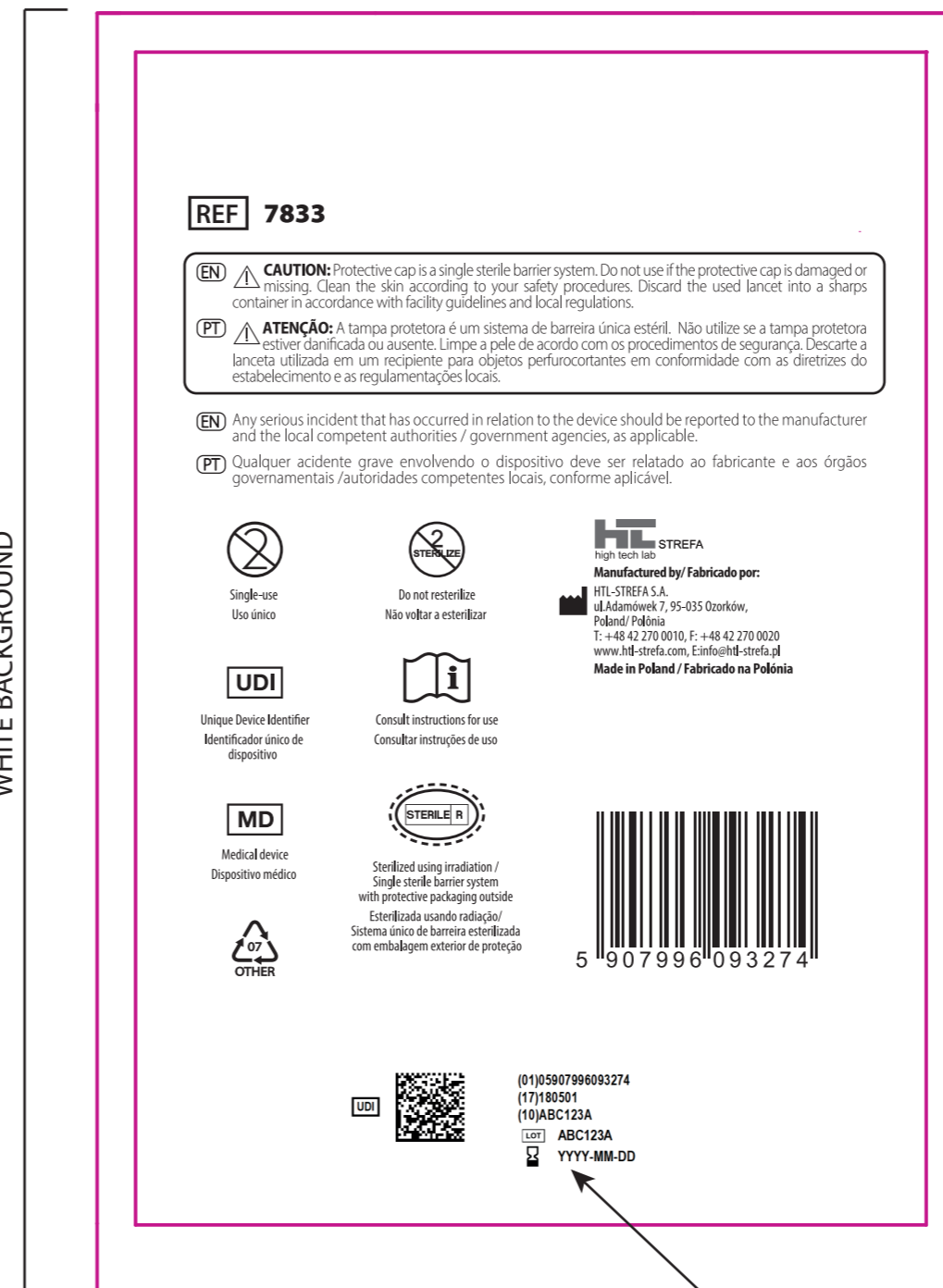
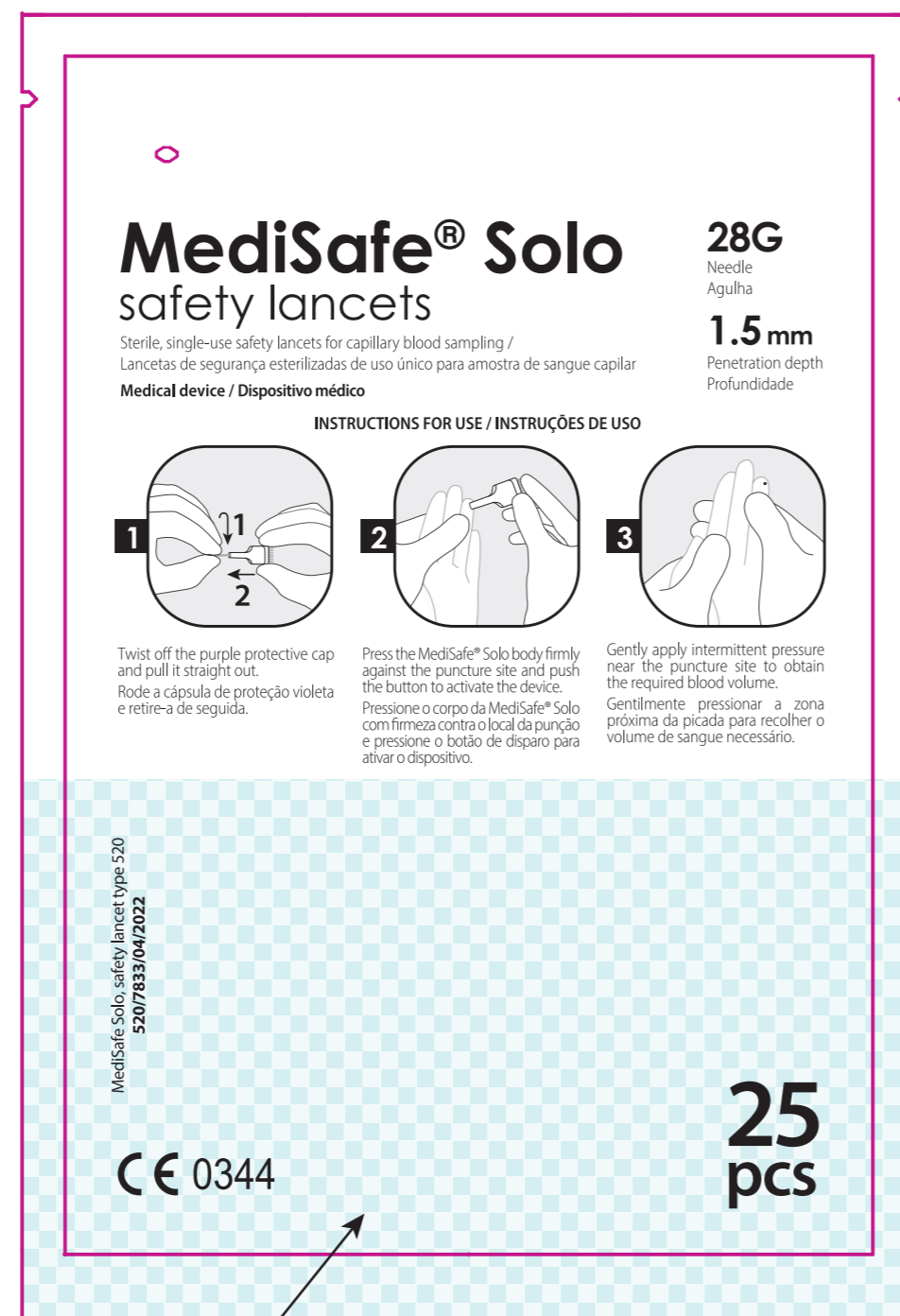
50mm

100mm

50mm

FRONT SIDE OF THE BAG

BACK SIDE OF THE BAG



- transparent window where the lancets are visible

UDI code printed directly on the bag during packing process

Instructions for use³

³ 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.

Abbott

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos de teste VIH-1/2**

Bioline™ HIV 1/2 3.0

The 3rd Generation of antibodies to HIV-1/HIV-2 Test

Le 3^e test de génération d'anticorps anti-VIH-1/2

O terceiro geração de anticorpos para a prueba de VIH 1 o 2

A 3.º **test de anticorpos do teste VIH-1/2**

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

Test procedure (Refer to figure)

1. Bring all kit components and specimens to reach a temperature between 15 °C and 30 °C prior to testing.
2. Remove the test device from foil pouch and place it on a flat, dry surface. Label the test device with patient identifier.
3. [Using a micropipette] Dispense 20 µl of plasma or serum specimen or dispense 20 µl of whole blood specimen into the specimen well.
OR,
[Using capillary pipette] Dispense 20 µl of drawn whole blood specimen into the specimen well.
4. Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well.
Caution: Do not let bottle nozzle touch device in order to avoid cross-contamination. Hold bottle vertically while dispensing. If you do not hold the bottle vertically, it can add inaccurate results. Exactly 4 drops should be added. Adding more than 4 drops may result in reddish color background or an invalid result.
5. As the test begins to work, you will see purple color move across the result window in the center of the test device.
6. Interpret test results 10 - 20 minutes after adding assay diluent. Do not read after 20 minutes.
Caution: If the test result is not legible after 10 minutes due to high background color, read again later but within 20 minutes of adding the diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

Test interpretation (Refer to figure)

1. A colored control line will appear at "C" in the result window to show that the test is working properly.
2. The "T" reaction of the result window indicates the test result.
Negative result: The presence of only the control line (C) within the result window indicates a negative HIV-1/2 result.
Positive result:
1) **HIV-1 positive:**
• The presence of both test line 1 (I) and the control line (C) indicates a positive result for anti-HIV-1.
• If the color intensity of the test line 1 darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
2) **HIV-2 positive:**
• The presence of both test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-2.
• If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
Both HIV-1 and HIV-2 Positive: The presence of the test line 1 (I), the test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.
Caution: Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homolog in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine your virus or diagnose a co-infection accurately, you must perform a confirmatory tests such as Western Blot test.

Materials provided and active ingredients of main components

1. The Bioline™ HIV 1/2 3.0 test kit contains the following items to perform the assay:
1-1. Bioline™ HIV 1/2 3.0 [30 tests/kit] Cat. No. 03FK10.
30 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL vial)
• 1 Instructions for use
1-2. Bioline™ HIV 1/2 3.0 [25 Tests/kit] Cat. No. 03FK16.
25 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL vial)
• 25 Capillary pipettes (20 µl), 25 Sterile lancets, 25 Alcohol swabs
• 1 Instructions for use
2. Active ingredients of main components
1. Test strip including Gold conjugate: Recombinant HIV-1 gp41, 24nt, HIV-2 gp36 antigen, gold colloid (0.050 ± 0.01 µg), Test line 1: Recombinant HIV-1 antigen (gp41, p24) (0.025±0.015 µg), Test line 2: Recombinant HIV-2 antigen (gp36) (0.05±0.01 µg), Control line (C) anti-HIV-1/2 IgG (0.5±0.05 µg)
2. Assay diluent: 50M Tris-HCl Buffer (pH 7.0), Sodium azide (0.4%)

Materials required but not provided

1. Micropipette, Protective gloves, Timer, Biohazard container

Kit storage and stability

1. The test kit should be stored at a temperature between 1 °C and 30 °C. Do not freeze the test kit components.
2. Assay diluent should be opened and resealed for each assay. Cap should be firmly sealed between each use. Assay diluent is stable until expiration date if kept at 1 - 30 °C.
3. The test device is sensitive to heat and humidity. Perform the test immediately after opening the test kit pouch.
4. Do not use the test kit beyond its expiration date. The shelf life of the kit is indicated on the outer package.
5. Do not use the test kit if the pouch is damaged or the seal is broken.

Warnings

1. The test devices are for *in vitro* diagnostic use only. Do not reuse the test device.
2. The instructions should be followed exactly as directed on the package. Any individual performing an assay with this product must be trained in its use and must be proficient.
3. Do not use the pipette with mouth, smoke, drink, eat, apply cosmetics, or handle contact lenses in areas where the pipette is used.
4. Wear protective gloves while handling specimens and wash hands thoroughly afterwards.
5. Clean up spills thoroughly using an appropriate disinfectant.
6. Decontaminate and dispose of all specimens, reaction kits and potentially contaminated materials in a biohazard container as if they were infectious waste.
7. Do not mix or interchange different specimens.
8. Do not use the test kit for the detection of HIV-1 and anti-HIV-2.
9. Avoid splashing or aerosol formation of specimen and assay diluent.
10. Do not mix or interchange components among different lots or those for other products.
11. Do not drink assay diluent.
12. Care should be taken to avoid contamination of the bottle nozzle when dropping assay diluent into the specimen well.
13. The assay diluent contains a proprietary antimicrobial agent, sodium azide, which prevents the growth of bacteria. Do not use assay diluent for normal laboratory safety precautions as follows: If contact with assay diluent to the eyes and/or skin, wash affected area with soap and water immediately. If irritation or signs of toxicity occur, seek medical attention.
14. The assay diluent contains sodium azide, which may react with lead or copper plating to form highly explosive metal azide compounds. When disposing of these reagents through plumbing fixtures, flush with a large volume of water to prevent azide build-up in drains.
15. Safety data sheet available for professional user on request.

Specimen collection and handling

1. **Whole blood**
[Collection by micropipette]
• Using venipuncture, collect whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate).
• The blood specimen should be refrigerated at 2 - 8 °C.
• If stored at 2 - 8 °C, the blood specimen must be tested within 3 days of refrigeration.
• Do not use a blood specimen stored for more than 3 days; it can cause a nonspecific reaction.
• Bring blood specimens to room temperature (15 - 30 °C) prior to use.
[Collection using a lancet]
• Clean the area to be lanced with an alcohol swab.
• Squeeze the fingertip then prick the side of the finger with a lancet provided. Wipe away the first blood drop (then, safely) disposed of the lancet immediately after use.
• Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.
2. **Plasma or serum**
[Plasma]
• Using venipuncture, draw the whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate) and then centrifuge blood to obtain the plasma specimen.
[Serum]
• Using venipuncture, draw the whole blood into the collection tube (NOT containing anticoagulants) then leave for 30 minutes to allow blood coagulation to occur. Centrifuge the tube to separate the serum and discard the clot.
• If plasma or serum specimens are not tested immediately, they must be refrigerated at 2 - 8 °C. For storage period longer than 2 weeks, freezing (below -20 °C) is required. Bring plasma or serum specimens to room temperature (15 - 30 °C) prior to use.
3. **Precautions**
• Plasma or serum specimens containing a precipitate may yield inaccurate test results. Such specimens must be clarified by centrifugation prior to assaying.
• Anticoagulants including heparin, EDTA and sodium citrate do not affect the test result.
• Use of other anticoagulants than those listed will not be validated. Their use may affect the test result.
• Use new pipette tips for each specimen in order to avoid cross-contamination of other specimens which could cause erroneous results.

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

Test procedure (Refer to figure)

1. Bring all kit components and specimens to reach a temperature between 15 °C and 30 °C prior to testing.
2. Remove the test device from foil pouch and place it on a flat, dry surface. Label the test device with patient identifier.
3. [Using a micropipette] Dispense 20 µl of plasma or serum specimen or dispense 20 µl of whole blood specimen into the specimen well.
OR,
[Using capillary pipette] Dispense 20 µl of drawn whole blood specimen into the specimen well.
4. Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well.
Caution: Do not let bottle nozzle touch device in order to avoid cross-contamination. Hold bottle vertically while dispensing. If you do not hold the bottle vertically, it can add inaccurate results. Exactly 4 drops should be added. Adding more than 4 drops may result in reddish color background or an invalid result.
5. As the test begins to work, you will see purple color move across the result window in the center of the test device.
6. Interpret test results 10 - 20 minutes after adding assay diluent. Do not read after 20 minutes.
Caution: If the test result is not legible after 10 minutes due to high background color, read again later but within 20 minutes of adding the diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

Test interpretation (Refer to figure)

1. A colored control line will appear at "C" in the result window to show that the test is working properly.
2. The "T" reaction of the result window indicates the test result.
Negative result: The presence of only the control line (C) within the result window indicates a negative HIV-1/2 result.
Positive result:
1) **HIV-1 positive:**
• The presence of both test line 1 (I) and the control line (C) indicates a positive result for anti-HIV-1.
• If the color intensity of the test line 1 darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
2) **HIV-2 positive:**
• The presence of both test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-2.
• If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
Both HIV-1 and HIV-2 Positive: The presence of the test line 1 (I), the test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.
Caution: Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homolog in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine your virus or diagnose a co-infection accurately, you must perform a confirmatory tests such as Western Blot test.

Materials provided and active ingredients of main components

1. The Bioline™ HIV 1/2 3.0 test kit contains the following items to perform the assay:
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1-2. Bioline™ HIV 1/2 3.0 [25 Tests/kit] Cat. No. 03FK16.
25 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL vial)
• 25 Capillary pipettes (20 µl), 25 Sterile lancets, 25 Alcohol swabs
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2. Active ingredients of main components
1. Test strip including Gold conjugate: Recombinant HIV-1 gp41, 24nt, HIV-2 gp36 antigen, gold colloid (0.050 ± 0.01 µg), Test line 1: Recombinant HIV-1 antigen (gp41, p24) (0.025±0.015 µg), Test line 2: Recombinant HIV-2 antigen (gp36) (0.05±0.01 µg), Control line (C) anti-HIV-1/2 IgG (0.5±0.05 µg)
2. Assay diluent: 50M Tris-HCl Buffer (pH 7.0), Sodium azide (0.4%)

Materials required but not provided

1. Micropipette, Protective gloves, Timer, Biohazard container

Kit storage and stability

1. The test kit should be stored at a temperature between 1 °C and 30 °C. Do not freeze the test kit components.
2. Assay diluent should be opened and resealed for each assay. Cap should be firmly sealed between each use. Assay diluent is stable until expiration date if kept at 1 - 30 °C.
3. The test device is sensitive to heat and humidity. Perform the test immediately after opening the test kit pouch.
4. Do not use the test kit beyond its expiration date. The shelf life of the kit is indicated on the outer package.
5. Do not use the test kit if the pouch is damaged or the seal is broken.

Warnings

1. The test devices are for *in vitro* diagnostic use only. Do not reuse the test device.
2. The instructions should be followed exactly as directed on the package. Any individual performing an assay with this product must be trained in its use and must be proficient.
3. Do not use the pipette with mouth, smoke, drink, eat, apply cosmetics, or handle contact lenses in areas where the pipette is used.
4. Wear protective gloves while handling specimens and wash hands thoroughly afterwards.
5. Clean up spills thoroughly using an appropriate disinfectant.
6. Decontaminate and dispose of all specimens, reaction kits and potentially contaminated materials in a biohazard container as if they were infectious waste.
7. Do not mix or interchange different specimens.
8. Do not use the test kit for the detection of HIV-1 and anti-HIV-2.
9. Avoid splashing or aerosol formation of specimen and assay diluent.
10. Do not mix or interchange components among different lots or those for other products.
11. Do not drink assay diluent.
12. Care should be taken to avoid contamination of the bottle nozzle when dropping assay diluent into the specimen well.
13. The assay diluent contains a proprietary antimicrobial agent, sodium azide, which prevents the growth of bacteria. Do not use assay diluent for normal laboratory safety precautions as follows: If contact with assay diluent to the eyes and/or skin, wash affected area with soap and water immediately. If irritation or signs of toxicity occur, seek medical attention.
14. The assay diluent contains sodium azide, which may react with lead or copper plating to form highly explosive metal azide compounds. When disposing of these reagents through plumbing fixtures, flush with a large volume of water to prevent azide build-up in drains.
15. Safety data sheet available for professional user on request.

Specimen collection and handling

1. **Whole blood**
[Collection by micropipette]
• Using venipuncture, collect whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate).
• The blood specimen should be refrigerated at 2 - 8 °C.
• If stored at 2 - 8 °C, the blood specimen must be tested within 3 days of refrigeration.
• Do not use a blood specimen stored for more than 3 days; it can cause a nonspecific reaction.
• Bring blood specimens to room temperature (15 - 30 °C) prior to use.
[Collection using a lancet]
• Clean the area to be lanced with an alcohol swab.
• Squeeze the fingertip then prick the side of the finger with a lancet provided. Wipe away the first blood drop (then, safely) disposed of the lancet immediately after use.
• Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.
2. **Plasma or serum**
[Plasma]
• Using venipuncture, draw the whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate) and then centrifuge blood to obtain the plasma specimen.
[Serum]
• Using venipuncture, draw the whole blood into the collection tube (NOT containing anticoagulants) then leave for 30 minutes to allow blood coagulation to occur. Centrifuge the tube to separate the serum and discard the clot.
• If plasma or serum specimens are not tested immediately, they must be refrigerated at 2 - 8 °C. For storage period longer than 2 weeks, freezing (below -20 °C) is required. Bring plasma or serum specimens to room temperature (15 - 30 °C) prior to use.
3. **Precautions**
• Plasma or serum specimens containing a precipitate may yield inaccurate test results. Such specimens must be clarified by centrifugation prior to assaying.
• Anticoagulants including heparin, EDTA and sodium citrate do not affect the test result.
• Use of other anticoagulants than those listed will not be validated. Their use may affect the test result.
• Use new pipette tips for each specimen in order to avoid cross-contamination of other specimens which could cause erroneous results.

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

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Test procedure (Refer to figure)

1. Bring all kit components and specimens to reach a temperature between 15 °C and 30 °C prior to testing.
2. Remove the test device from foil pouch and place it on a flat, dry surface. Label the test device with patient identifier.
3. [Using a micropipette] Dispense 20 µl of plasma or serum specimen or dispense 20 µl of whole blood specimen into the specimen well.
OR,
[Using capillary pipette] Dispense 20 µl of drawn whole blood specimen into the specimen well.
4. Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well.
Caution: Do not let bottle nozzle touch device in order to avoid cross-contamination. Hold bottle vertically while dispensing. If you do not hold the bottle vertically, it can add inaccurate results. Exactly 4 drops should be added. Adding more than 4 drops may result in reddish color background or an invalid result.
5. As the test begins to work, you will see purple color move across the result window in the center of the test device.
6. Interpret test results 10 - 20 minutes after adding assay diluent. Do not read after 20 minutes.
Caution: If the test result is not legible after 10 minutes due to high background color, read again later but within 20 minutes of adding the diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

Test interpretation (Refer to figure)

1. A colored control line will appear at "C" in the result window to show that the test is working properly.
2. The "T" reaction of the result window indicates the test result.
Negative result: The presence of only the control line (C) within the result window indicates a negative HIV-1/2 result.
Positive result:
1) **HIV-1 positive:**
• The presence of both test line 1 (I) and the control line (C) indicates a positive result for anti-HIV-1.
• If the color intensity of the test line 1 darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
2) **HIV-2 positive:**
• The presence of both test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-2.
• If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
Both HIV-1 and HIV-2 Positive: The presence of the test line 1 (I), the test line 2 (II) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.
Caution: Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homolog in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine your virus or diagnose a co-infection accurately, you must perform a confirmatory tests such as Western Blot test.

Materials provided and active ingredients of main components

1. The Bioline™ HIV 1/2 3.0 test kit contains the following items to perform the assay:
1-1. Bioline™ HIV 1/2 3.0 [30 tests/kit] Cat. No. 03FK10.
30 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL vial)
• 1 Instructions for use
1-2. Bioline™ HIV 1/2 3.0 [25 Tests/kit] Cat. No. 03FK16.
25 Test devices with desiccant in individual foil pouches
• Assay diluent (1 x 4 mL vial)
• 25 Capillary pipettes (20 µl), 25 Sterile lancets, 25 Alcohol swabs
• 1 Instructions for use
2. Active ingredients of main components
1. Test strip including Gold conjugate: Recombinant HIV-1 gp41, 24nt, HIV-2 gp36 antigen, gold colloid (0.050 ± 0.01 µg), Test line 1: Recombinant HIV-1 antigen (gp41, p24) (0.025±0.015 µg), Test line 2: Recombinant HIV-2 antigen (gp36) (0.05±0.01 µg), Control line (C) anti-HIV-1/2 IgG (0.5±0.05 µg)
2. Assay diluent: 50M Tris-HCl Buffer (pH 7.0), Sodium azide (0.4%)

Materials required but not provided

1. Micropipette, Protective gloves, Timer, Biohazard container

Kit storage and stability

1. The test kit should be stored at a temperature between 1 °C and 30 °C. Do not freeze the test kit components.
2. Assay diluent should be opened and resealed for each assay. Cap should be firmly sealed between each use. Assay diluent is stable until expiration date if kept at 1 - 30 °C.
3. The test device is sensitive to heat and humidity. Perform the test immediately after opening the test kit pouch.
4. Do not use the test kit beyond its expiration date. The shelf life of the kit is indicated on the outer package.
5. Do not use the test kit if the pouch is damaged or the seal is broken.

Warnings

1. The test devices are for *in vitro* diagnostic use only. Do not reuse the test device.
2. The instructions should be followed exactly as directed on the package. Any individual performing an assay with this product must be trained in its use and must be proficient.
3. Do not use the pipette with mouth, smoke, drink, eat, apply cosmetics, or handle contact lenses in areas where the pipette is used.
4. Wear protective gloves while handling specimens and wash hands thoroughly afterwards.
5. Clean up spills thoroughly using an appropriate disinfectant.
6. Decontaminate and dispose of all specimens, reaction kits and potentially contaminated materials in a biohazard container as if they were infectious waste.
7. Do not mix or interchange different specimens.
8. Do not use the test kit for the detection of HIV-1 and anti-HIV-2.
9. Avoid splashing or aerosol formation of specimen and assay diluent.
10. Do not mix or interchange components among different lots or those for other products.
11. Do not drink assay diluent.
12. Care should be taken to avoid contamination of the bottle nozzle when dropping assay diluent into the specimen well.
13. The assay diluent contains a proprietary antimicrobial agent, sodium azide, which prevents the growth of bacteria. Do not use assay diluent for normal laboratory safety precautions as follows: If contact with assay diluent to the eyes and/or skin, wash affected area with soap and water immediately. If irritation or signs of toxicity occur, seek medical attention.
14. The assay diluent contains sodium azide, which may react with lead or copper plating to form highly explosive metal azide compounds. When disposing of these reagents through plumbing fixtures, flush with a large volume of water to prevent azide build-up in drains.
15. Safety data sheet available for professional user on request.

Specimen collection and handling

1. **Whole blood**
[Collection by micropipette]
• Using venipuncture, collect whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate).
• The blood specimen should be refrigerated at 2 - 8 °C.
• If stored at 2 - 8 °C, the blood specimen must be tested within 3 days of refrigeration.
• Do not use a blood specimen stored for more than 3 days; it can cause a nonspecific reaction.
• Bring blood specimens to room temperature (15 - 30 °C) prior to use.
[Collection using a lancet]
• Clean the area to be lanced with an alcohol swab.
• Squeeze the fingertip then prick the side of the finger with a lancet provided. Wipe away the first blood drop (then, safely) disposed of the lancet immediately after use.
• Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.
2. **Plasma or serum**
[Plasma]
• Using venipuncture, draw the whole blood into the collection tube (containing anticoagulant including heparin, EDTA and sodium citrate) and then centrifuge blood to obtain the plasma specimen.
[Serum]
• Using venipuncture, draw the whole blood into the collection tube (NOT containing anticoagulants) then leave for 30 minutes to allow blood coagulation to occur. Centrifuge the tube to separate the serum and discard the clot.
• If plasma or serum specimens are not tested immediately, they must be refrigerated at 2 - 8 °C. For storage period longer than 2 weeks, freezing (below -20 °C) is required. Bring plasma or serum specimens to room temperature (15 - 30 °C) prior to use.
3. **Precautions**
• Plasma or serum specimens containing a precipitate may yield inaccurate test results. Such specimens must be clarified by centrifugation prior to assaying.
• Anticoagulants including heparin, EDTA and sodium citrate do not affect the test result.
• Use of other anticoagulants than those listed will not be validated. Their use may affect the test result.
• Use new pipette tips for each specimen in order to avoid cross-contamination of other specimens which could cause erroneous results.

El **3.º** **test de generación de anticorpos anti-VIH-1/2** es el tercer generacón de anticorpos para la prueba de VIH 1 o 2. A 3.º **test de anticorpos do teste VIH-1/2**

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FRANÇAIS

À propos du test
[Précautions] Le VIH (virus de l'immunodéficience humaine) est reconnu comme l'agent étiologique du syndrome d'immunodéficience acquise (SIDA). El virus se transmet par contacts sexuels, exposition à du sang infecté, à certains liquides ou tissus corporels, et de la mère au fœtus ou à l'enfant pendant le période périnatale. Le VIH-1 est isolé à partir de patients atteints du SIDA et du complexe liés à SIDA, et de personnes en bonne santé ayant un potentiel élevé de risque de développer le SIDA. Les patients infectés par le VIH-2 se trouvent principalement dans des régions d'Afrique d'Ouest. Son évolution est marquée par deux niveaux croissants de réplication virale et émergence de souches virales plus virulentes. Le VIH-1 et le VIH-2 sont semblables en ce qui concerne leur morphologie, tropisme cellulaire, interaction avec l'hôte et structure générique. Des études sérologiques ont déterminé que le VIH-1 et le VIH-2 ont plusieurs antigènes sérologiques communs dans le sang total humain par immunodéficience acquise. Ce diagnostic clinique du VIH peut inclure la détection d'anticorpos contre le VIH-1 ou le VIH-2 en plasma ou le sérum humain par immunodéficience. La présence du VIH peut être identifiée par la détection d'anticorpos sérologiques communs dans le sang total humain par immunodéficience. Ce diagnostic clinique du VIH peut inclure la détection d'anticorpos anti-VIH 1/2 dans le plasma ou le sérum humain par immunodéficience. La présence du VIH peut être identifiée par la détection d'anticorpos sérologiques communs dans le sang total humain par immunodéficience. Ce diagnostic clinique du VIH peut inclure la détection d'anticorpos anti-VIH 1/2 dans le plasma ou le sérum humain par immunodéficience. La présence du VIH peut être identifiée par la détection d'anticorpos sérologiques communs dans le sang total humain par immunodéficience.

- Utilisez un nouvel embout de pipette pour chaque échantillon afin d'éviter toute contamination croisée d'autres échantillons qui pourrait entraîner des résultats erronés.
- Ne réutilisez pas les composants du kit et les échantillons à une température comprise entre 15 et 30 °C avant d'effectuer le test.
- Retirez le dispositif de test de la pochette en aluminium et placez-le sur une surface plane et sèche. Apposez sur le dispositif de test une étiquette comportant l'identifiant du patient. [À l'aide d'une micropipette]
 - Versez 10 µl d'échantillon de plasma ou de sérum ou 20 µl d'échantillon de sang total dans le puits d'échantillonnage + S.
 - [À l'aide d'une pipette capillaire] Versez 20 µl d'échantillon de sang total prélevé dans le puits d'échantillonnage + S.
 - Versez 4 gouttes (environ 120 µl) de diluente dans le puits d'échantillonnage + S.
 - Attention:** Veillez à ce que l'embout du flacon ne touche pas le dispositif afin d'éviter toute contamination croisée. Tenez le flacon à la verticale pour éviter fesserchillement. Vous ne tenez pas le flacon à la verticale, des résultats inexactes risquent d'être obtenus. Vous devez attendre 4 gouttes exactement. L'appât de plus de 4 gouttes peut donner lieu à un arrière-plan de couleur rougeâtre ou à un résultat non valide.
 - Lorsque le test commence à faire effet, une couleur violette se déplace à travers la fenêtre de résultat au centre du dispositif de test.
 - Interprétez les résultats du test au bout de 10 à 20 minutes après l'ajout du diluant de dosage. Ne lisez pas les résultats après 20 minutes.
 - Attention:** Le kit Bioline™ HIV 1/2 3.0 est un test qualitatif rapide pour la détection d'anticorpos de tous les isotopes (IgG, IgM et IgA) spécifiques au VIH-1 et au VIH-2 simultanément dans le sérum, le plasma ou le sang total humain. Le kit Bioline™ HIV 1/2 3.0 est destiné uniquement à un usage diagnostique et à diagnostic *in vitro*. Ce test peut ne pas convenir au diagnostic d'infection précoce ni au dépistage lors d'un don de sang. Les échantillons positifs doivent être confirmés par un dosage supplémentaire tel que le test ELISA ou le test Western Blot. Les performances du dosage n'ont pas été établies pour les populations de nourissons, d'enfants ou d'adultes.

Interprétation du test (voir les illustrations)

1. Une ligne de contrôle colorée apparaît au niveau de la lettre « C », dans la fenêtre de résultat, pour indiquer que le test fonctionne correctement.
2. La partie « T » et 2 de la fenêtre de résultat indique le résultat du test.
Résultat négatif: La présence, dans la fenêtre de résultat, de la ligne de contrôle (C) uniquement indique un résultat négatif.

- Matériel fourni et substances actives des principaux composants**
1. Le kit de test Bioline™ HIV 1/2 3.0 comporte les éléments suivants pour la réalisation du dosage:
1-1. Bioline™ HIV 1/2 3.0 [30 tests/kit] N. cat. 03FK10.
30 Dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium
• Diluant de test (1 x 4 mL/flacon)
• 1 Mode d'emploi
1-2. Bioline™ HIV 1/2 3.0 [25 tests/kit] N. cat. 03FK16.
25 Dispositifs de test avec agent déshydratant conditionnés dans des emballages individuels en aluminium
• Diluant de test (1 x 4 mL/flacon)
• 25 pipettes capillaires (20 µl), 25 lancettes stériles, 25 compresses d'alcool
• 1 Mode d'emploi
2. Ingrédients actifs des principaux composants
1) **Positif anti-VIH-1:**
• La présence simultanée de la ligne de test 1 (I) et de la ligne de contrôle (C) indique un résultat positif pour les anticorpos anti-VIH-2.
• Si l'intensité de couleur de la ligne de test 1 est plus sombre que celle de la ligne de test 2 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorpos anti-VIH-1.
2) **Positif anti-VIH-2:**
• La présence simultanée de la ligne de test 2 (II) et de la ligne de contrôle (C) indique un résultat positif pour les anticorpos anti-VIH-2.
• Si l'intensité de couleur de la ligne de test 2 est plus sombre que celle de la ligne de test 1 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorpos anti-VIH-2.
3) **Positif aux deux VIH-1 et VIH-2:** La présence de la ligne de test 1 (I), de la ligne de test 2 (II) et de la ligne de contrôle (C) indique un résultat positif pour les anticorpos anti-VIH-1 et/ou anti-VIH-2.
Attention: Un patient positif result positif pour les anticorps anti-VIH-1 et anti-VIH-2, ce qui est habitent son cas en rare, il est possible qu'il existe une homologie dans la séquence d'acides aminés entre anticorps anti-VIH-1 et anti-VIH-2. Pour déterminer le type de virus ou diagnostiquer une co-infection avec précision, vous devez effectuer un test de confirmation, par exemple le test Western Blot.
Résultat non valide:
• Si l'intensité de couleur de la ligne de test 1 est plus sombre que celle de la ligne de test 2 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorpos anti-VIH-1.
• Si l'intensité de couleur de la ligne de test 2 est plus sombre que celle de la ligne de test 1 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorps anti-VIH-2.
Positif aux deux VIH-1 et VIH-2: La présence de la ligne de test 1 (I), de la ligne de test 2 (II) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-1 et/ou anti-VIH-2.
Attention: Un patient positif result positif pour les anticorps anti-VIH-1 et anti-VIH-2, ce qui est habitent son cas en rare, il est possible qu'il existe une homologie dans la séquence d'acides aminés entre anticorps anti-VIH-1 et anti-VIH-2. Pour déterminer le type de virus ou diagnostiquer une co-infection avec précision, vous devez effectuer un test de confirmation, par exemple le test Western Blot.

Conservation et stabilité du kit

- Le kit de test doit être conservé à une température comprise entre 1 et 30 °C. Ne congélez pas les composants.
- Le diluant de dosage doit être remis en état et refermé à chaque dosage. Veillez à bien refermer le flacon à l'aide du bouchon entre chaque utilisation. S'il est conservé à une température comprise entre 1 et 30 °C, le diluant de dosage est stable jusqu'à la date de péremption.
- Le dispositif de test est stable à température ambiante. Effectuez le test immédiatement après avoir retiré le dispositif de test de la pochette en aluminium.
- N'utilisez pas le kit de test au-delà de sa date de péremption. La durée de conservation du kit est indiquée sur l'emballage extérieur.
- N'utilisez pas le kit de test si la pochette est endommagée ou si le sceau est rompu.

Avertissements

- Les dispositifs de test sont uniquement destinés au diagnostic *in vitro*. Ne réutilisez pas le kit de test.
- Pour obtenir des résultats exacts et précis, il est impératif de suivre scrupuleusement les instructions fournies. Toute personne réalisant un dosage à l'aide de ce produit doit avoir été formée à son utilisation et être compétente en la matière.
- N'utilisez pas la pipette avec le bouche,

Bioline[®] HIV 1/2 3.0

The 3rd Generation of antibodies to HIV-1/HIV-2 Test
Le test de 3^e génération d'anticorps anti-VIH-1/VIH-2
La tercera generación de anticuerpos para la prueba de VIH 1 o 2
A 3.^a geração de anticorpos do teste VIH-1/VIH-2

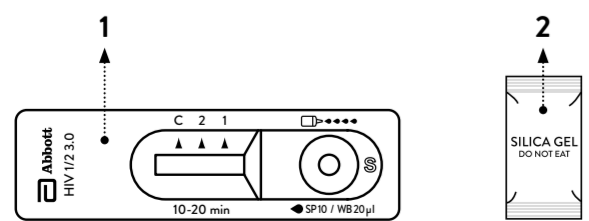
PREPARATION / PRÉPARATION / PREPARACIÓN / PREPARAÇÃO

- 1** **EN** Now, open the package and look for the following:
1. Test device with desiccant in individual foil pouch
2. Assay diluent
3. Instructions for use
- FR** Ouvrez l'emballage et vérifiez que les éléments suivants sont présents :
1. Dispositif de test avec agent déshydratant conditionné dans un emballage en aluminium individuel
2. Diluant du test
3. Mode d'emploi
- ES** Ahora, abra el paquete y busque lo siguiente:
1. Dispositivo de prueba con desecante en la bolsa de papel de aluminio individual
2. Diluyente del ensayo
3. Instrucciones de uso
- PT** Agora, abra a embalagem e procure o seguinte:
1. Dispositivo de teste com dessecante em bolsa de alumínio individual
2. Diluente de ensaio
3. Instruções de utilização
- Including only for Catalog No. 03FK16**
4. Capillary pipette (20 µl)
5. Sterile lancet
6. Alcohol swab
- Inclus uniquement pour le N° de cat. 03FK16**
4. Pipette capillaire (20 µl)
5. Lancette stérile
6. Compresse d'alcool
- Lo siguiente solo se incluye para el n.º de catálogo 03FK16**
4. Pipeta capilar (20 µl)
5. Lanceta estéril
6. Hisopo con alcohol

- 2** **EN** Carefully read the instructions on how to use the Bioline™ HIV 1/2 3.0 test kit.
- FR** Lisez attentivement les instructions d'utilisation du kit de test Bioline™ HIV 1/2 3.0.
- ES** Lea atentamente las instrucciones sobre cómo utilizar el kit de prueba Bioline™ HIV 1/2 3.0.
- PT** Leia atentamente as instruções sobre como utilizar o kit de teste Bioline™ HIV 1/2 3.0.

- 3** **EN** Look at the expiration date at the back of the foil pouch. If the expiration date has passed, use another kit. To avoid false results, ensure that the assay diluent used is from the same kit as the new test device.
- FR** Vérifiez la date de péremption au dos de la pochette en aluminium. Si la date de péremption est dépassée, utilisez un autre kit. Pour éviter l'obtention de résultats erronés, assurez-vous que le diluant utilisé pour le dosage provient du même kit que le nouveau dispositif de test.
- ES** Revise la fecha de vencimiento en la parte posterior de la bolsa de papel aluminio. Si el kit está vencido, utilice otro kit. Para evitar resultados falsos, asegúrese de que el diluyente del análisis que se utilice sea del mismo kit que el dispositivo de prueba nuevo.
- PT** Verifique a data de validade na parte posterior da bolsa de alumínio. Se a data de validade tiver expirado, utilize outro kit. Para evitar resultados falsos, certifique-se de que o diluente de ensaio utilizado é do mesmo kit que o novo dispositivo de teste.

- 4** **EN** Open the foil pouch and look for the following:
1. Test device
2. Desiccant
Then, label the device with the patient identifier.
- FR** Ouvrez la pochette en aluminium et vérifiez la présence des éléments suivants :
1. Dispositif de test
2. Dessiccant
Apposez ensuite sur le dispositif une étiquette comportant l'identifiant du patient.
- ES** Abra la bolsa de papel aluminio y busque lo siguiente:
1. Dispositivo de prueba
2. Desecante
A continuación, etiquete el dispositivo con el identificador del paciente.
- PT** Abra a bolsa de alumínio e procure o seguinte:
1. Dispositivo de teste
2. Dessecante
Em seguida, coloque uma etiqueta no dispositivo com o identificador do paciente.



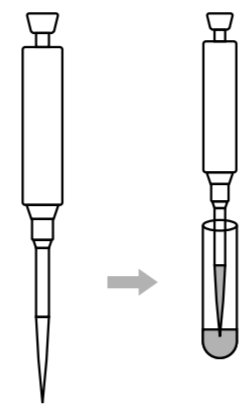
◀ SP10 µl / WB20 µl : Serum 10 µl or Plasma 10 µl or Whole blood 20 µl / Sérum 10 µl ou plasma 10 µl ou sang total 20 µl / 10 µl de soro, 10 µl de plasma o 20 µl de sangue / 10 µl de soro, 10 µl de plasma ou 20 µl de sangue total
▶ ◀◀◀◀◀ Assay diluent 4 drops / 4 gouttes de diluant de dosage / 4 gotas de diluyente del análisis / 4 gotas de diluente de ensaio

TEST PROCEDURE / DÉROULEMENT DU TEST / PROCEDIMIENTO DE PRUEBA / PROCEDIMENTO DO TESTE

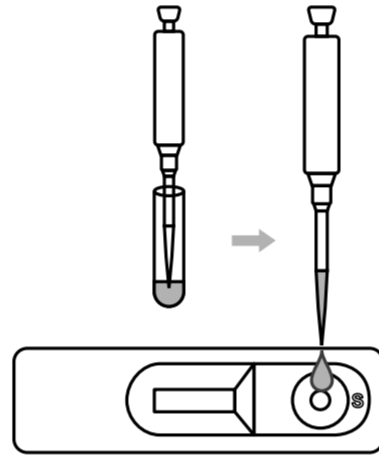
I. Blood (by venipuncture), Plasma or Serum specimen / Échantillon de sang (par ponction veineuse), de plasma ou de sérum / Muestra de sangre (por venopunción), plasma o suero / Amostra de sangue (por punção venosa), plasma ou soro

Specimen collection / Prélèvement d'un échantillon / Obtención de la muestra / Colheita de amostras

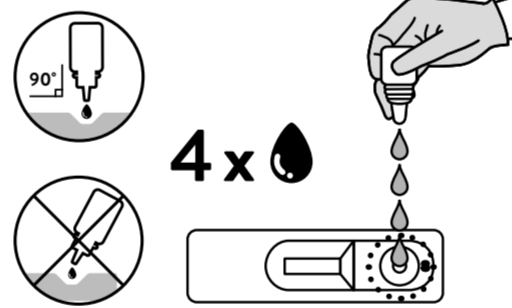
- 1** **EN** Using a micropipette draw plasma or serum: 10 µl specimen or whole blood: 20 µl specimen
- FR** À l'aide d'une micropipette, prélevez du plasma ou du sérum : échantillon de 10 µl ou sang total : échantillon de 20 µl
- ES** Con una micropipeta, extraiga plasma o suero: muestra de 10 µl o sangre: muestra de 20 µl
- PT** Utilizando uma micropipeta, colha plasma ou soro: amostra de 10 µl ou sangue total: amostra de 20 µl



- 1** **EN** Dispense 10 µl of plasma or serum specimen or 20 µl of whole blood specimen into the specimen well marked "S"
- FR** Versez 10 µl d'échantillon de plasma ou de sérum ou 20 µl d'échantillon de sang total dans le puits d'échantillonnage marqué « S »
- ES** Vierta 10 µl de la muestra de plasma o de suero o 20 µl de la muestra de sangre en el pocillo para muestra marcado con una "S"
- PT** Coloque 10 µl da amostra de plasma ou soro ou 20 µl da amostra de sangue total no poço da amostra com a marca "S"



- 2** **EN** Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.
- FR** Versez 4 gouttes (environ 120 µl) de diluant de dosage dans le puits d'échantillonnage « S ». Tenez le flacon à la verticale pour verser l'échantillon. Veillez à ce que l'embout du flacon ne touche pas le dispositif afin d'éviter toute contamination croisée.
- ES** Vierta 4 gotas (aproximadamente 120 µl) de diluyente del análisis en el pocillo para prueba "S". Sostenga la botella en posición vertical mientras vierte. Tenga cuidado de que la boquilla de la botella no entre en contacto con el dispositivo para evitar la contaminación cruzada.
- PT** Coloque 4 gotas (aproximadamente, 120 µl) de diluente de ensaio no poço da amostra "S". Segure o frasco na vertical durante a colocação das gotas. Não deixe que o bocal do frasco toque no dispositivo para evitar a contaminação cruzada.



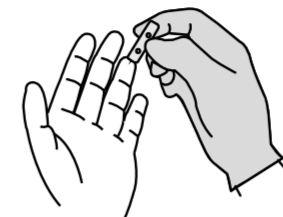
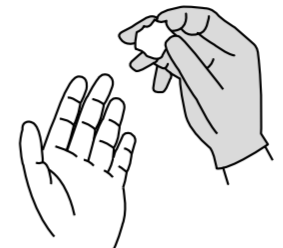
- 3** **EN** Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.
- FR** Interprétez les résultats au bout de 10 à 20 minutes après l'ajout du diluant de dosage. Une lecture en dehors de cette période (avant 10 min ou après 20 min) peut donner lieu à des résultats erronés.
- ES** Interprete el resultado de 10 a 20 minutos después de agregar el diluyente del análisis. La lectura fuera de este marco de tiempo (antes de 10 min o después de 20 min) puede arrojar resultados falsos.
- PT** Interprete o resultado 10 a 20 minutos depois de adicionar o diluente de ensaio. A leitura fora deste intervalo de tempo (antes de 10 min ou após 20 min) pode fornecer resultados falsos.



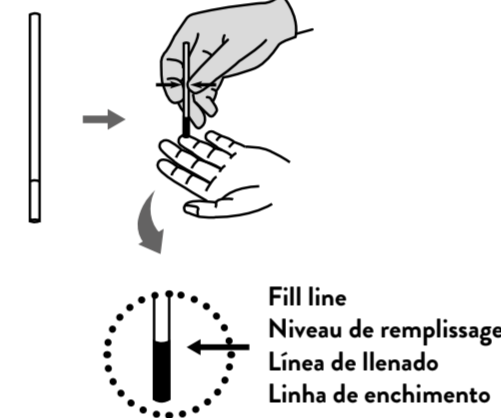
II. Blood specimen (with a lancet) / Échantillon de sang (prélevé à l'aide d'une lancette) / Muestra de sangre (con lanceta) / Amostra de sangue (com uma lanceta)

Specimen collection / Prélèvement d'un échantillon / Obtención de la muestra / Colheita de amostras

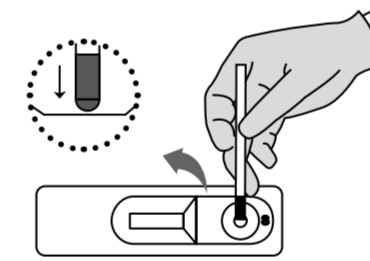
- 1** **EN** Clean the area to be lanced with an alcohol swab.
FR Nettoyez la zone de prélèvement avec un tampon imbibé d'alcool.
ES Limpie el área desde donde se va a extraer la sangre con un hisopo con alcohol.
PT Limpe a área a lancetar, utilizando uma zaragatoa com álcool.
- 2** **EN** Squeeze the fingertip then prick the lateral side of the finger with a lancet provided. Wipe away the first blood drop. Then, safely dispose of the lancet immediately after.
FR Appuyez fermement sur le bout du doigt, puis piquez sur le côté du doigt avec la lancette fournie. Essuyez la première goutte de sang. Mettez immédiatement la lancette au rebut selon la procédure de sécurité prévue.
ES Apriete la punta del dedo y, a continuación, realice una punción en la parte lateral del dedo con una lanceta proporcionada. Limpie la primera gota de sangre. Inmediatamente después, deseché la lanceta de forma segura.
PT Aperte a ponta do dedo e, em seguida, pique a parte lateral do dedo com a lanceta fornecida. Limpe a primeira gota de sangue. Imediatamente depois, elimine a lanceta num recipiente apropriado.



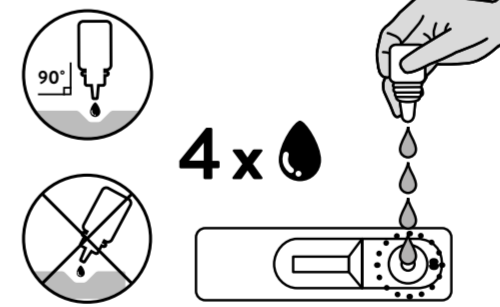
- 3** **EN** Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.
- FR** Plongez l'extrémité ouverte d'une nouvelle pipette capillaire (20 µl) dans la goutte de sang suivante et relâchez la pression pour que la pipette capillaire aspire le sang jusqu'à atteindre le niveau de remplissage.
- ES** Sumerja el extremo abierto de una pipeta capilar nueva (20 µl) en la siguiente gota de sangre que aflore y libere la presión, para que entre sangre en la pipeta capilar hasta la línea de llenado.
- PT** Mergulhe a extremidade aberta de uma nova pipeta capilar (20 µl) na próxima gota de sangue e liberte a pressão para colher o sangue para a pipeta capilar até à linha de enchimento.



- 1** **EN** Dispense 20 µl of drawn whole blood specimen in the specimen well marked "S".
FR Versez 20 µl d'échantillon de sang total prélevé dans le puits d'échantillonnage marqué « S ».
ES Vierta 20 µl de la muestra de sangre extraída en el pocillo para muestra marcado con una "S".
PT Coloque 20 µl da amostra de sangue total colhido no poço da amostra com a marca "S".



- 2** **EN** Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.
FR Versez 4 gouttes (environ 120 µl) de diluant de dosage dans le puits d'échantillonnage « S ». Tenez le flacon à la verticale pour verser l'échantillon. Veillez à ce que l'embout du flacon ne touche pas le dispositif afin d'éviter toute contamination croisée.
ES Vierta 4 gotas (aproximadamente 120 µl) de diluyente del análisis en el pocillo para prueba "S". Sostenga la botella en posición vertical mientras vierte. Tenga cuidado de que la boquilla de la botella no entre en contacto con el dispositivo para evitar la contaminación cruzada.
PT Coloque 4 gotas (aproximadamente, 120 µl) de diluente de ensaio no poço da amostra "S". Segure o frasco na vertical durante a colocação das gotas. Não deixe que o bocal do frasco toque no dispositivo para evitar a contaminação cruzada.



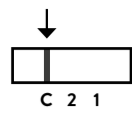
- 3** **EN** Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.
FR Interprétez les résultats au bout de 10 à 20 minutes après l'ajout du diluant de dosage. Une lecture en dehors de cette période (avant 10 min ou après 20 min) peut donner lieu à des résultats erronés.
ES Interprete el resultado de 10 a 20 minutos después de agregar el diluyente del análisis. La lectura fuera de este marco de tiempo (antes de 10 min o después de 20 min) puede arrojar resultados falsos.
PT Interprete o resultado 10 a 20 minutos depois de adicionar o diluente de ensaio. A leitura fora deste intervalo de tempo (antes de 10 min ou após 20 min) pode fornecer resultados falsos.



INTERPRETATION / INTERPRÉTATION / INTERPRETACIÓN / INTERPRETAÇÃO

Negative / Négatif / Negativo

- EN** The presence of only the control line (C) within the result window indicates a negative result.
FR La présence, dans la fenêtre de résultat, de la ligne de contrôle (C) uniquement indique un résultat négatif.
ES Si solo aparece la línea de control (C) en la ventana de resultados, significa que el resultado de la prueba es negativo.
PT A presença de apenas a linha de controlo (C) na janela de resultados indica um resultado negativo.

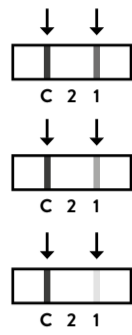


Positive / Positif / Positivo

- EN** ⚠ **Caution:** The presence of any test line, no matter how faint, the result is considered positive.
FR ⚠ **Attention :** la présence d'une ligne de test, quelle que soit l'intensité de la couleur, indique un résultat considéré comme positif.
ES ⚠ **Precaución:** Si aparece cualquier línea de prueba, sin importar lo tenue que sea, significa que el resultado de la prueba es positivo.
PT ⚠ **Atenção:** a presença de qualquer linha de teste, independentemente da intensidade desta, significa que o resultado é considerado positivo.

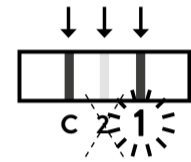
HIV-1 Positive / Positif / Positivo

- 2 LINES / 2 LIGNES / 2 LÉNES / 2 LINHAS**
- EN** The presence of both test line 1 (1) and the control line (C) indicates a positive result for anti-HIV-1.
FR La présence simultanée de la ligne de test 1 (1) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-1.
ES La presencia de las líneas de prueba 1 (1) y de control (C) indica un resultado positivo para anticuerpos anti-VIH-1.
PT A presença da linha de teste 1 (1) e da linha de controlo (C) indica um resultado positivo para anti-VIH-1.
- Strong / Forts / Intensa / Forte
- Medium / Moyens / Mediana / Médio
- Weak / Faibles / Tenue / Ténue



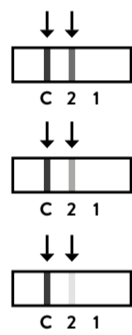
3 LINES / 3 LIGNES / 3 LÉNES / 3 LINHAS

- EN** If the color intensity of the test line 1 is darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.
FR Si l'intensité de couleur de la ligne de test 1 est plus sombre que celle de la ligne de test 2 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorps anti-VIH-1.
Ligne pâle : par homologie dans la séquence d'acides aminés du VIH de type 1 et du VIH de type 2.
ES Si la intensidad del color de la línea de prueba 1 es más oscura que la de la línea de prueba 2 en la ventana de resultados, se puede interpretar como un resultado positivo para anticuerpos anti-VIH-1.
Línea borrosa: por homología en la secuencia de aminoácidos de VIH Tipo 1 y 2.
PT Se, na janela de resultados, a intensidade da cor da linha de teste 1 for mais escura do que a da linha de teste 2, pode interpretar o resultado como positivo para anti-VIH-1.
Linha tênue: por homologia na sequência de aminoácidos do VIH tipo 1 e VIH tipo 2.



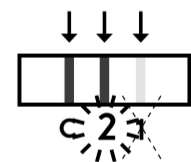
HIV-2 Positive / Positif / Positivo

- 2 LINES / 2 LIGNES / 2 LÉNES / 2 LINHAS**
- EN** The presence of both test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-2.
FR La présence simultanée de la ligne de test 2 (2) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-2.
ES La presencia de las líneas de prueba 2 (2) y de control (C) indica un resultado positivo para anticuerpos anti-VIH-2.
PT A presença da linha de teste 2 (2) e da linha de controlo (C) indica um resultado positivo para anti-VIH-2.
- Strong / Forts / Intensa / Forte
- Medium / Moyens / Mediana / Médio
- Weak / Faibles / Tenue / Ténue



3 LINES / 3 LIGNES / 3 LÉNES / 3 LINHAS

- EN** If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.
FR Si l'intensité de couleur de la ligne de test 2 est plus sombre que celle de la ligne de test 1 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorps anti-VIH-2.
Ligne pâle : par homologie dans la séquence d'acides aminés du VIH de type 1 et du VIH de type 2.
ES Si la intensidad del color de la línea de prueba 2 es más oscura que la de la línea de prueba 1 en la ventana de resultados, se puede interpretar como un resultado positivo para anticuerpos anti-VIH-2.
Línea borrosa: por homología en la secuencia de aminoácidos de VIH Tipo 1 y 2.
PT Se, na janela de resultados, a intensidade da cor da linha de teste 2 for mais escura do que a da linha de teste 1, pode interpretar o resultado como positivo para anti-VIH-2.
Linha tênue: por homologia na sequência de aminoácidos do VIH tipo 1 e VIH tipo 2.

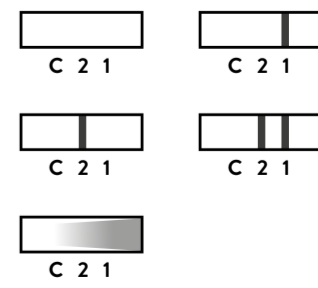


Remark / Remarque / Comentarios / Observação

- EN** [Both HIV-1 and HIV-2 Positive]
The presence of the test line 1 (1), the test line 2 (2) and the control line (C) indicates a positive result for anti-VIH-1 and/or anti-VIH-2.
⚠ **Caution:** Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homology in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine the virus type or diagnose a co-infection accurately, you must perform a confirmatory test such as Western Blot etc.
- FR** [Positifs aux deux VIH-1 et VIH-2]
La présence de la ligne de test 1 (1), de la ligne de test 2 (2) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-1 et/ou anti-VIH-2.
⚠ **Attention :** bien qu'un résultat positif pour les anticorps anti-VIH-1 et anti-VIH-2 chez un patient soit un cas rare, il est possible qu'il existe une homologie dans la séquence d'acides aminés entre les anticorps anti-VIH-1 et anti-VIH-2. Pour déterminer le type de virus ou diagnostiquer une co-infection avec précision, vous devez effectuer un test de confirmation, tel que le test Western Blot, etc.
- ES** [Resultado positivo para VIH 1 y 2]
La presencia de las líneas de prueba 1 (1), de prueba 2 (2) y de control (C) indica un resultado positivo para anticuerpos anti-VIH 1 y 2.
⚠ **Precaución:** Si bien es poco común que un paciente presente un resultado positivo para ambos anticuerpos, anti-VIH 1 y 2, esto está dentro de las posibilidades, debido a que la secuencia de aminoácidos entre anticuerpos anti-VIH 1 y 2 es homóloga. Para determinar el tipo de virus o diagnosticar una coinfección con precisión, se debe realizar una prueba de confirmación, como Western Blot, etc.
- PT** [Positivo para VIH-1 e VIH-2]
A presença da linha de teste 1 (1), da linha de teste 2 (2) e da linha de controlo (C) indica um resultado positivo para anticorpos anti-VIH-1 e/ou anti-VIH-2.
⚠ **Atenção:** apesar de ser um caso raro, é possível obter um resultado positivo para o anti-VIH-1 e o anti-VIH-2 num único paciente, pois existe homologia na sequência de aminoácidos entre o anti-VIH-1 e o anti-VIH-2. Para determinar o tipo de vírus ou diagnosticar uma coinfeção com precisão, é necessário realizar um teste de confirmação, tal como o Western Blot, etc.

Invalid / Non valide / No válido / Inválido

- EN** Absence of the control line (C) and/or presence of a pink/purple smear in the result window indicates an invalid result. Instructions may not have been followed correctly or the test may have deteriorated. It is recommended that the specimen be retested using a new test device.
- FR** L'absence de la ligne de contrôle (C) et/ou la présence d'un frottis rose/violet dans la fenêtre de résultat indiquent un résultat non valide. Il est possible que les instructions n'aient pas été suivies correctement ou que le test se soit détérioré. Il est recommandé de tester à nouveau l'échantillon à l'aide d'un nouveau dispositif de test.
- ES** Si no aparece la línea de control (C) o si aparece una mancha rosada o púrpura en la ventana de resultados, significa que el resultado de la prueba no es válido. Es posible que no se hayan seguido las instrucciones correctamente o que la prueba se haya deteriorado. Se recomienda que la muestra se vuelva a analizar con un nuevo dispositivo de prueba.
- PT** A ausência da linha de controlo (C) e/ou a presença de uma mancha rosada ou púrpura na janela de resultados indica um resultado inválido. As instruções podem não ter sido seguidas corretamente ou o teste pode ter-se deteriorado. Recomenda-se que a amostra seja novamente testada com um novo dispositivo de teste.



Bioline™ HIV 1/2 3.0

The 3rd Generation of antibodies to HIV-1/HIV-2 Test
 Le test de 3e génération d'anticorps anti-VIH-1/VIH-2
 La tercera generación de anticuerpos para la prueba de VIH 1 o 2
 A 3.ª geração de anticorpos do teste VIH-1/VIH-2

PREPARATION / PRÉPARATION / PREPARACIÓN / PREPARAÇÃO

- EN Now, open the package and look for the following:

 - Test device with desiccant in individual foil pouch
 - Assay diluent
 - Instructions for use
 - Capillary pipette (20 µl)
 - Safety lancet
 - Alcohol swab

ES Ahora, abra el paquete y busque lo siguiente:

 - Dispositivo de prueba con desecante en la bolsa de papel de aluminio individual
 - Diluyente del ensayo
 - Instrucciones de uso
 - Pipeta capilar (20 µl)
 - Lancetas de seguridad
 - Hisopo con alcohol

FR Ouvrez l'emballage et vérifiez que les éléments suivants sont présents :

 - Dispositif de test avec agent déshydratant conditionné dans un emballage en aluminium individuel
 - Diluant du test
 - Mode d'emploi
 - Pipette capillaire (20 µl)
 - Lancettes de sécurité
 - Compresse d'alcool

PT Agora, abra a embalagem e procure o seguinte:

 - Dispositivo de teste com dessecante em bolsa de alumínio individual
 - Diluyente de ensaio
 - Instruções de utilização
 - Pipeta capilar (20 µl)
 - Lancetas de segurança
 - Zaragatoa com álcool

- EN Carefully read the instructions on how to use the Bioline™ HIV 1/2 3.0 test kit.

ES Lea atentamente las instrucciones sobre cómo utilizar el kit de prueba Bioline™ HIV 1/2 3.0.

FR Lisez attentivement les instructions d'utilisation du kit de test Bioline™ HIV 1/2 3.0.

PT Leia atentamente as instruções sobre como utilizar o kit de teste Bioline™ HIV 1/2 3.0.

- EN Look at the expiration date at the back of the foil pouch. If the expiration date has passed, use another kit. To avoid false results, ensure that the assay diluent used is from the same kit as the new test device.

FR Vérifiez la date de péremption au dos de la pochette en aluminium. Si la date de péremption est dépassée, utilisez un autre kit. Pour éviter l'obtention de résultats erronés, assurez-vous que le diluant utilisé pour le dosage provient du même kit que le nouveau dispositif de test.

ES Revise la fecha de vencimiento en la parte posterior de la bolsa de papel aluminio. Si el kit está vencido, utilice otro kit. Para evitar resultados falsos, asegúrese de que el diluyente del análisis que se utilice sea del mismo kit que el dispositivo de prueba nuevo.

PT Verifique a data de validade na parte posterior da bolsa de alumínio. Se a data de validade tiver expirado, utilize outro kit. Para evitar resultados falsos, certifique-se de que o diluente de ensaio utilizado é do mesmo kit que o novo dispositivo de teste.

- EN Open the foil pouch and look for the following:

 - Test device
 - Desiccant

Then, label the device with the patient identifier.

FR Ouvrez la pochette en aluminium et vérifiez la présence des éléments suivants :

 - Dispositif de test
 - Desiccant

Apposez ensuite sur le dispositif une étiquette comportant l'identifiant du patient.

ES Abra la bolsa de papel aluminio y busque lo siguiente:

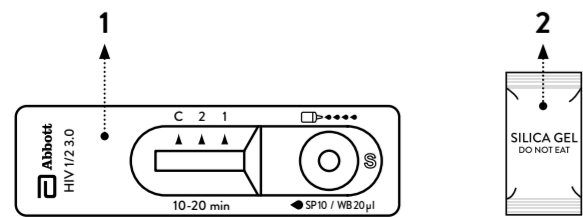
 - Dispositivo de prueba
 - Desecante

A continuación, etiquete el dispositivo con el identificador del paciente.

PT Abra a bolsa de alumínio e procure o seguinte:

 - Dispositivo de teste
 - Dessecante

Em seguida, coloque uma etiqueta no dispositivo com o identificador do paciente.



● SP 10µl / WB 20µl / Serum 10 µl or Plasma 10 µl or Whole blood 20 µl / Sérum 10 µl ou plasma 10 µl ou sang total 20 µl / 10 µl de soro, 10 µl de plasma o 20 µl de sangue / 10 µl de soro, 10 µl de plasma ou 20 µl de sangue total
 ● Assay diluent 4 drops / 4 gouttes de diluant de dosage / 4 gotas de diluyente del análisis / 4 gotas de diluente de ensaio

TEST PROCEDURE / DÉROULEMENT DU TEST / PROCEDIMIENTO DE PRUEBA / PROCEDIMENTO DO TESTE

I. Blood (by venipuncture), Plasma or Serum specimen / Échantillon de sang (par ponction veineuse), de plasma ou de sérum / Muestra de sangre (por venopunción), plasma o suero / Amostra de sangue (por punção venosa), plasma ou soro

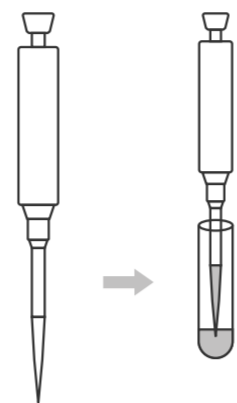
Specimen collection / Prélèvement d'un échantillon / Obtención de la muestra / Colheita de amostras

- EN Using a micropipette draw plasma or serum: 10 µl specimen or whole blood: 20 µl specimen

FR À l'aide d'une micropipette, prélevez du plasma ou du sérum : échantillon de 10 µl ou sang total : échantillon de 20 µl

ES Con una micropipeta, extraiga plasma o suero: muestra de 10 µl o sangre: muestra de 20 µl

PT Utilizando uma micropipeta, colha plasma ou soro: amostra de 10 µl ou sangue total: amostra de 20 µl

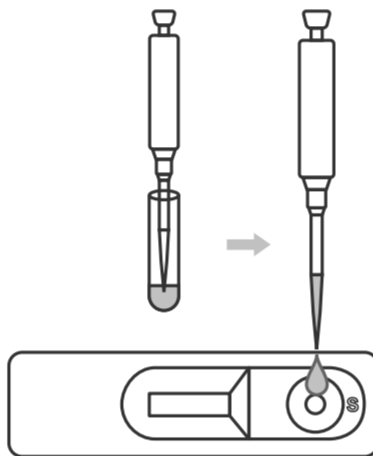


- EN Dispense 10 µl of plasma or serum specimen or 20 µl of whole blood specimen into the specimen well marked "S"

FR Versez 10 µl d'échantillon de plasma ou de sérum ou 20 µl d'échantillon de sang total dans le puits d'échantillonnage marqué « S »

ES Vierta 10 µl de la muestra de plasma o de suero o 20 µl de la muestra de sangre en el pocillo para muestra marcado con una "S"

PT Coloque 10 µl da amostra de plasma ou soro ou 20 µl da amostra de sangue total no poço da amostra com a marca "S"

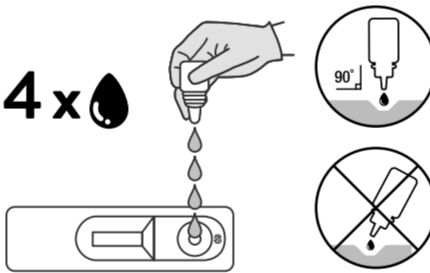


- EN Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.

FR Versez 4 gouttes (environ 120 µl) de diluant de dosage dans le puits d'échantillonnage « S ». Tenez le flacon à la verticale pour verser l'échantillon. Veillez à ce que l'embout du flacon ne touche pas le dispositif afin d'éviter toute contamination croisée.

ES Vierta 4 gotas (aproximadamente 120 µl) de diluyente del análisis en el pocillo para prueba "S". Sostenga la botella en posición vertical mientras vierte. Tenga cuidado de que la boquilla de la botella no entre en contacto con el dispositivo para evitar la contaminación cruzada.

PT Coloque 4 gotas (aproximadamente, 120 µl) de diluente de ensaio no poço da amostra "S". Segure o frasco na vertical durante a colocação das gotas. Não deixe que o bocal do frasco toque no dispositivo para evitar a contaminação cruzada.



- EN Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

FR Interprétez les résultats au bout de 10 à 20 minutes après l'ajout du diluant de dosage. Une lecture en dehors de cette période (avant 10 min ou après 20 min) peut donner lieu à des résultats erronés.

ES Interprete el resultado de 10 a 20 minutos después de agregar el diluyente del análisis. La lectura fuera de este marco de tiempo (antes de 10 min o después de 20 min) puede arrojar resultados falsos.

PT Interprete o resultado 10 a 20 minutos depois de adicionar o diluente de ensaio. A leitura fora deste intervalo de tempo (antes de 10 min ou após 20 min) pode fornecer resultados falsos.



II. Blood specimen (with a lancet) / Échantillon de sang (prélevé à l'aide d'une lancette) / Muestra de sangre (con lanceta) / Amostra de sangue (com uma lanceta)

Specimen collection / Prélèvement d'un échantillon / Obtención de la muestra / Colheita de amostras

- EN Clean the area to be lanced with an alcohol swab.

FR Nettoyez la zone de prélèvement avec un tampon imbibé d'alcool.

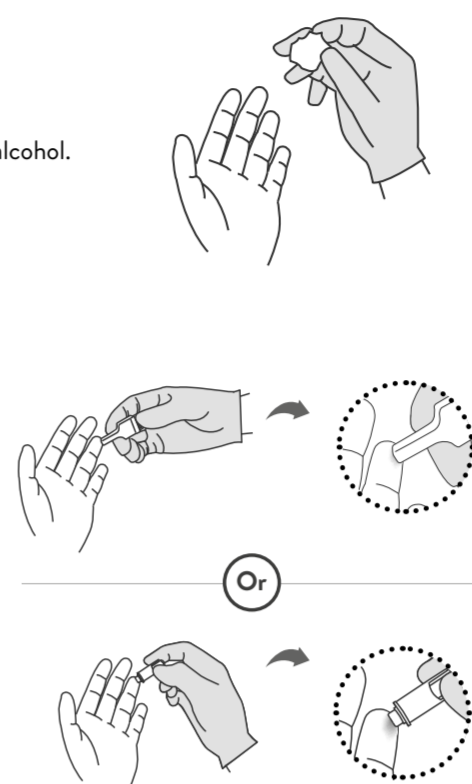
ES Limpie el área desde donde se va a extraer la sangre con un hisopo con alcohol.

PT Limpe a área a lancetar, utilizando uma zaragatoa com álcool.
- EN Squeeze the fingertip then prick the lateral side of the finger with a lancet provided. Wipe away the first blood drop. Then, safely dispose of the lancet immediately after.

FR Appuyez fermement sur le bout du doigt, puis piquez sur le côté du doigt avec la lancette fournie. Essuyez la première goutte de sang. Mettez immédiatement la lancette au rebut selon la procédure de sécurité prévue.

ES Apriete la punta del dedo y, a continuación, realice una punción en la parte lateral del dedo con una lanceta proporcionada. Limpie la primera gota de sangre. Inmediatamente después, deseche la lanceta de forma segura.

PT Aperte a ponta do dedo e, em seguida, pique a parte lateral do dedo com a lanceta fornecida. Limpe a primeira gota de sangue. Imediatamente depois, elimine a lanceta num recipiente apropriado.

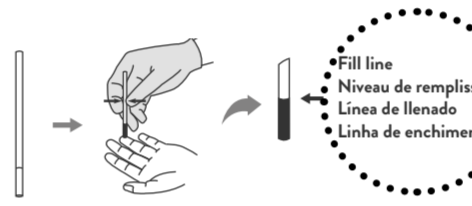


- EN Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.

FR Plongez l'extrémité ouverte d'une nouvelle pipette capillaire (20 µl) dans la goutte de sang suivante et relâchez la pression pour que la pipette capillaire aspire le sang jusqu'à atteindre le niveau de remplissage.

ES Sumerja el extremo abierto de una pipeta capilar nueva (20 µl) en la siguiente gota de sangre que aflore y libere la presión, para que entre sangre en la pipeta capilar hasta la línea de llenado.

PT Mergulhe a extremidade aberta de uma nova pipeta capilar (20 µl) na próxima gota de sangue e liberte a pressão para colher o sangue para a pipeta capilar até à linha de enchimento.



- EN Dispense 20 µl of drawn whole blood specimen in the specimen well marked "S".

FR Versez 20 µl d'échantillon de sang total prélevé dans le puits d'échantillonnage marqué « S ».

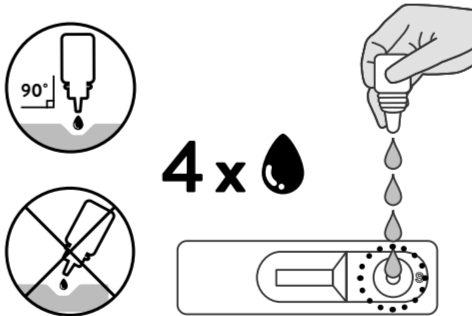
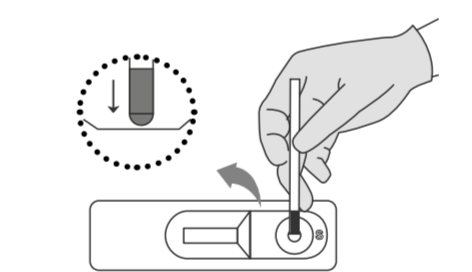
ES Vierta 20 µl de la muestra de sangre extraída en el pocillo para muestra marcado con una "S".

PT Coloque 20 µl da amostra de sangue total colhido no poço da amostra com a marca "S".
- EN Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.

FR Versez 4 gouttes (environ 120 µl) de diluant de dosage dans le puits d'échantillonnage « S ». Tenez le flacon à la verticale pour verser l'échantillon. Veillez à ce que l'embout du flacon ne touche pas le dispositif afin d'éviter toute contamination croisée.

ES Vierta 4 gotas (aproximadamente 120 µl) de diluyente del análisis en el pocillo para prueba "S". Sostenga la botella en posición vertical mientras vierte. Tenga cuidado de que la boquilla de la botella no entre en contacto con el dispositivo para evitar la contaminación cruzada.

PT Coloque 4 gotas (aproximadamente, 120 µl) de diluente de ensaio no poço da amostra "S". Segure o frasco na vertical durante a colocação das gotas. Não deixe que o bocal do frasco toque no dispositivo para evitar a contaminação cruzada.



- EN Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

FR Interprétez les résultats au bout de 10 à 20 minutes après l'ajout du diluant de dosage. Une lecture en dehors de cette période (avant 10 min ou après 20 min) peut donner lieu à des résultats erronés.

ES Interprete el resultado de 10 a 20 minutos después de agregar el diluyente del análisis. La lectura fuera de este marco de tiempo (antes de 10 min o después de 20 min) puede arrojar resultados falsos.

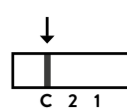
PT Interprete o resultado 10 a 20 minutos depois de adicionar o diluente de ensaio. A leitura fora deste intervalo de tempo (antes de 10 min ou após 20 min) pode fornecer resultados falsos.



INTERPRETATION / INTERPRÉTATION / INTERPRETAÇÃO

Negative / Négatif / Negativo

- EN The presence of only the control line (C) within the result window indicates a negative result.
- FR La présence, dans la fenêtre de résultat, de la ligne de contrôle (C) uniquement indique un résultat négatif.
- ES Si solo aparece la línea de control (C) en la ventana de resultados, significa que el resultado de la prueba es negativo.
- PT A presença de apenas a linha de controlo (C) na janela de resultados indica um resultado negativo.

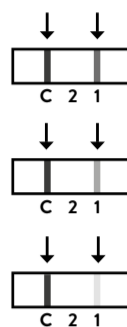


Positive / Positif / Positivo

- EN ⚠ Caution: The presence of any test line, no matter how faint, the result is considered positive.
- FR ⚠ Attention : la présence d'une ligne de test, quelle que soit l'intensité de la couleur, indique un résultat considéré comme positif.
- ES ⚠ Precaución: Si aparece cualquier línea de prueba, sin importar lo tenue que sea, significa que el resultado de la prueba es positivo.
- PT ⚠ Atenção: a presença de qualquer linha de teste, independentemente da intensidade desta, significa que o resultado é considerado positivo.

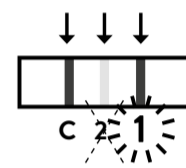
HIV-1 Positive / Positif / Positivo

- 2 LINES / 2 LIGNES / 2 LÍNEAS / 2 LINHAS**
- EN The presence of both test line 1 (1) and the control line (C) indicates a positive result for anti-HIV-1.
- FR La présence simultanée de la ligne de test 1 (1) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-1.
- ES La presencia de las líneas de prueba 1 (1) y de control (C) indica un resultado positivo para anticuerpos anti-VIH-1.
- PT A presença da linha de teste 1 (1) e da linha de controlo (C) indica um resultado positivo para anti-VIH-1.
- Strong / Forts / Intensa / Forte
- Medium / Moyens / Mediana / Médio
- Weak / Faibles / Tenue / Ténue



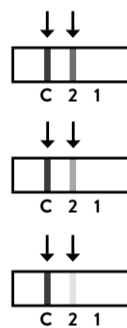
3 LINES / 3 LIGNES / 3 LÍNEAS / 3 LINHAS

- EN If the color intensity of the test line 1 is darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
- FR Si l'intensité de couleur de la ligne de test 1 est plus sombre que celle de la ligne de test 2 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorps anti-VIH-1.
- ES Si la intensidad del color de la línea de prueba 1 es más oscura que la de la línea de prueba 2 en la ventana de resultados, se puede interpretar como un resultado positivo para anticuerpos anti-VIH-1.
- PT Se, na janela de resultados, a intensidade da cor da linha de teste 1 for mais escura do que a da linha de teste 2, pode interpretar o resultado como positivo para anti-VIH-1.
- Linea borrosa: por homologia en la secuencia de aminoácidos de VIH Tipo 1 y 2.
- EN Ligne pâle : par homologie dans la séquence d'acides aminés du VIH de type 1 et du VIH de type 2.
- FR Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.
- ES Linea borrosa: por homología en la secuencia de aminoácidos de VIH Tipo 1 y 2.
- PT Se, na janela de resultados, a intensidade da cor da linha de teste 1 for mais escura do que a da linha de teste 2, pode interpretar o resultado como positivo para anti-VIH-1.
- Linea tênue: por homologia na sequência de aminoácidos do VIH tipo 1 e VIH tipo 2.



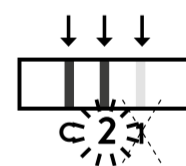
HIV-2 Positive / Positif / Positivo

- 2 LINES / 2 LIGNES / 2 LÍNEAS / 2 LINHAS**
- EN The presence of both test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-2.
- FR La présence simultanée de la ligne de test 2 (2) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-2.
- ES La presencia de las líneas de prueba 2 (2) y de control (C) indica un resultado positivo para anticuerpos anti-VIH-2.
- PT A presença da linha de teste 2 (2) e da linha de controlo (C) indica um resultado positivo para anti-VIH-2.
- Strong / Forts / Intensa / Forte
- Medium / Moyens / Mediana / Médio
- Weak / Faibles / Tenue / Ténue



3 LINES / 3 LIGNES / 3 LÍNEAS / 3 LINHAS

- EN If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
- FR Si l'intensité de couleur de la ligne de test 2 est plus sombre que celle de la ligne de test 1 dans la fenêtre de résultat, vous pouvez interpréter le résultat comme étant positif aux anticorps anti-VIH-2.
- ES Si la intensidad del color de la línea de prueba 2 es más oscura que la de la línea de prueba 1 en la ventana de resultados, se puede interpretar como un resultado positivo para anticuerpos anti-VIH-2.
- PT Se, na janela de resultados, a intensidade da cor da linha de teste 2 for mais escura do que a da linha de teste 1, pode interpretar o resultado como positivo para anti-VIH-2.
- Linea borrosa: por homologia en la secuencia de aminoácidos de VIH Tipo 1 y 2.
- EN Ligne pâle : par homologie dans la séquence d'acides aminés du VIH de type 1 et du VIH de type 2.
- FR Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.
- ES Linea borrosa: por homología en la secuencia de aminoácidos de VIH Tipo 1 y 2.
- PT Se, na janela de resultados, a intensidade da cor da linha de teste 2 for mais escura do que a da linha de teste 1, pode interpretar o resultado como positivo para anti-VIH-2.
- Linea tênue: por homologia na sequência de aminoácidos do VIH tipo 1 e VIH tipo 2.

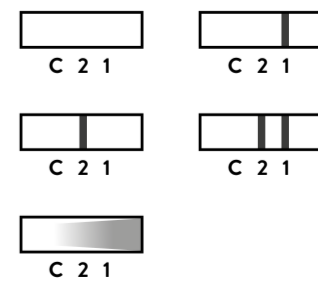


Remark / Remarque / Comentarios / Observação

- EN [Both HIV-1 and HIV-2 Positive] The presence of the test line 1 (1), the test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.
- FR [Positifs aux deux VIH-1 et VIH-2] La présence de la ligne de test 1 (1), de la ligne de test 2 (2) et de la ligne de contrôle (C) indique un résultat positif pour les anticorps anti-VIH-1 et/ou anti-VIH-2.
- ES [Resultado positivo para VIH 1 y 2] La presencia de las líneas de prueba 1 (1), de prueba 2 (2) y de control (C) indica un resultado positivo para anticuerpos anti-VIH 1 y 2.
- PT [Positivo para VIH-1 e VIH-2] A presença da linha de teste 1 (1), da linha de teste 2 (2) e da linha de controlo (C) indica um resultado positivo para anticuerpos anti-VIH 1 y 2.
- EN ⚠ Caution: Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homology in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine the virus type or diagnose a co-infection accurately, you must perform a confirmatory test such as Western Blot etc.
- FR ⚠ Attention : bien qu'un résultat positif pour les anticorps anti-VIH-1 et anti-VIH-2 chez un patient soit un cas rare, il est possible qu'il existe une homologie dans la séquence d'acides aminés entre les anticorps anti-VIH-1 et anti-VIH-2. Pour déterminer le type de virus ou diagnostiquer une co-infection avec précision, vous devez effectuer un test de confirmation, tel que le test Western Blot, etc.
- ES ⚠ Precaución: Si bien es poco común que un paciente presente un resultado positivo para ambos anticuerpos, anti-VIH 1 y 2, esto está dentro de las posibilidades, debido a que la secuencia de aminoácidos entre anticuerpos anti-VIH-1 y 2 es homóloga. Para determinar el tipo de virus o diagnosticar una coinfección con precisión, se debe realizar una prueba de confirmación, como Western Blot, etc.
- PT ⚠ Atenção: apesar de ser um caso raro, é possível obter um resultado positivo para o anti-VIH-1 e o anti-VIH-2 num único paciente, pois existe homologia na sequência de aminoácidos entre o anti-VIH-1 e o anti-VIH-2. Para determinar o tipo de vírus ou diagnosticar uma coinfeção com precisão, é necessário realizar um teste de confirmação, tal como o Western Blot, etc.

Invalid / Non valide / No válido / Inválido

- EN Absence of the control line (C) and/or presence of a pink/purple smear in the result window indicates an invalid result. Instructions may not have been followed correctly or the test may have deteriorated. It is recommended that the specimen be retested using a new test device.
- FR L'absence de la ligne de contrôle (C) et/ou la présence d'un frottis rose/violet dans la fenêtre de résultat indiquent un résultat non valide. Il est possible que les instructions n'aient pas été suivies correctement ou que le test se soit détérioré. Il est recommandé de tester à nouveau l'échantillon à l'aide d'un nouveau dispositif de test.
- ES Si no aparece la línea de control (C) o si aparece una mancha rosada o púrpura en la ventana de resultados, significa que el resultado de la prueba no es válido. Es posible que no se hayan seguido las instrucciones correctamente o que la prueba se haya deteriorado. Se recomienda que la muestra se vuelva a analizar con un nuevo dispositivo de prueba.
- PT Se não aparece a linha de controlo (C) ou se aparece uma mancha rosada ou púrpura na janela de resultados, significa que o resultado de a prova não é válido. É possível que não se tenham seguido as instruções correctamente o que a prova se tenha deteriorado. Recomenda-se que a amostra seja novamente testada com um novo dispositivo de teste.





REF 03FK22

Bioline™ HIV 1/2 3.0

The 3rd Generation of antibodies to HIV-1/HIV-2 Test

About the test

[Introduction] HIV (Human Immunodeficiency Virus) is recognized as the etiologic agent of Acquired Immune Deficiency Syndrome (AIDS). The virus is transmitted by sexual contact, exposure to infected blood, certain body fluids or tissues, and from mother to fetus or child during the perinatal period. HIV-1 has been isolated from patients with AIDS and AIDS related complex, and from healthy persons with high potential risk of developing AIDS. Patients with HIV-2 are found primarily in parts of West Africa. Its course is marked by increasing levels of viral replication and the emergence of more virulent viral strains. HIV-1 and HIV-2 are similar in their morphology, cell tropism, host interaction and generic structure. Serological studies have determined that HIV-1 and HIV-2 have multiple common epitopes in core antigens but much less so in the envelope antigens. This clinical diagnosis of HIV may include the detection of antibodies to HIV 1/2 in human plasma or serum by immunoassay. The presence of HIV can be identified by detection of antibodies to HIV 1/2 in human serum, plasma and whole blood by immunoassay. This advanced assay utilizes recombinant antigens targeted against immunogenic proteins. The major immunoreactive antigens of these proteins are HIV-1 gp41, p24 and HIV-2 gp36.

[Test principle] The Bioline™ HIV 1/2 3.0 test contains a membrane strip, which is precoated with recombinant HIV-1 capture antigen (gp41, p24) on test line 1 region and with recombinant HIV-2 capture antigen (gp36) on test line 2 region respectively. The recombinant HIV 1/2 antigen (gp41, p24 and gp36)-colloid gold conjugate and the sample move along the membrane chromatographically to the test region (T) and forms a visible line as the antigen-antibody-antigen gold particle complex forms with high degree of sensitivity and specificity.

[Intended use] The Bioline™ HIV 1/2 3.0 kit is a rapid, qualitative test for the detection of antibodies to all isotypes (IgG, IgM, IgA) specific to HIV-1 and HIV-2 simultaneously in human serum, plasma or whole blood. The Bioline™ HIV 1/2 3.0 kit is intended only for professional use and for *in vitro* diagnostic use. This test may not be suitable for diagnosis of early infection or blood donation screening. Positive samples should be confirmed by a supplemental assay such as ELISA or Western Blot test. The performance of the assay has not been established for populations of infants, children or adults.

Materials provided and active ingredients of main components

- The Bioline™ HIV 1/2 3.0 test kit contains the following items to perform the assay:
 - 25 Test devices with desiccant in individual foil pouches
 - Assay diluent (5 x 4 ml/vial)
 - 25 Capillary pipettes (20 µl), 25 Sterile lancets, 25 Alcohol swabs
 - 5 Instructions for use
- Active ingredients of main components
 - 1 test strip included: Gold conjugate: Recombinant HIV-1 gp41, p24, HIV-2 gp36 antigen – gold colloid (1.0±0.2 µg), Test line 1: Recombinant HIV -1 antigen (gp41, p24) (0.625±0.125 µg), Test line 2: Recombinant HIV -2 antigen (gp36) (0.5±0.1 µg), Control line: Goat anti-HIV serum (0.75±0.15 µg)
 - Assay diluent: 50mM Tris-HCl Buffer (4 ml), Sodium azide (q.s.)

Materials required but not provided

- Micropipette, Protective gloves, Timer, Biohazard container

Kit storage and stability

- The test kit should be stored at a temperature between 1 °C and 30 °C. Do not freeze the kit or its components.
- Assay diluent may be opened and resealed for each assay. Cap should be firmly sealed between each use. Assay diluent is stable until expiration date if kept at 1 - 30 °C.
- The test device is sensitive to both heat and humidity. Perform the test immediately after removing the test device from foil pouch.
- Do not use the test kit beyond its expiration date. The shelf life of the kit is as indicated on the outer package.
- Do not use the test kit if the pouch is damaged or the seal is broken.

Warnings

- The test devices are for *in vitro* diagnostic use only. Do not reuse the test device.
- The instructions must be followed exactly to achieve accurate results. Any individual performing an assay with this product must be trained in its use and must be proficient.
- Do not use the pipette by mouth, smoke, drink, eat, apply cosmetics, or handle contact lenses in areas where specimens or kit components are being handled.
- Wear protective gloves while handling specimens and wash hands thoroughly afterwards.
- Clean up spills thoroughly using an appropriate disinfectant.
- Decontaminate and dispose of all specimens, reaction kits and potentially contaminated materials in a biohazard container as if they were infectious waste.
- Do not mix or interchange different specimens.
- Do not eat the desiccant from the foil pouch.
- Avoid splashing or aerosol formation of specimen and assay diluent.
- Do not mix or interchange components among different lots or those for other products.
- Do not drink assay diluent.
- Care should be taken to avoid contamination of the bottle nozzle when dropping assay diluent into the specimen well.
- The assay diluent contains a proprietary antimicrobial agent, sodium azide, which presents no hazard to the user if normal laboratory safety precautions are followed. If contact with assay diluent to the eyes and/or skin, wash affected area with soap and water immediately. If irritation or signs of toxicity occur, seek medical attention.
- The assay diluent contains sodium azide, which may react with lead or copper plumbing to form highly explosive metal azide compounds. When disposing of these reagents through plumbing fixtures, flush with a large volume of water to prevent azide build-up in drains.
- Safety data sheet available for professional user on request.

Specimen collection and handling

- Whole blood**
[Collection by venipuncture]
 - Using venipuncture, collect whole blood into the collection tube (containing anticoagulants including heparin, EDTA and sodium citrate).
 - If the blood specimen is not immediately tested, it must be refrigerated at 2 - 8 °C.
 - If stored at 2 - 8 °C, the blood specimen must be tested within 3 days of refrigeration.
 - Do not use a blood specimen stored for more than 3 days; it can cause a nonspecific reaction.
 - Bring blood specimens to room temperature (15 - 30 °C) prior to use.[Collection using a lancet]
 - Clean the area to be lanced with an alcohol swab.
 - Squeeze the fingertip then prick the lateral side of the finger with a lancet provided. Wipe away the first blood drop. Then, safely dispose of the lancet immediately after.
 - Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.
- Plasma or serum**
 - [Plasma] Using venipuncture, draw the whole blood into the collection tube (containing anticoagulants including heparin, EDTA and sodium citrate) and then centrifuge blood to obtain the plasma specimen.
 - [Serum] Using venipuncture, draw the whole blood into the collection tube (NOT containing anticoagulants) then leave for 30 minutes to allow blood coagulation to occur. Centrifuge the tube to generate a serum specimen.
 - If plasma or serum specimens are not tested immediately, they must be refrigerated at 2 - 8 °C. For storage period longer than 2 weeks, freezing (below -20 °C) is required. Bring plasma or serum specimens to room temperature (15 - 30 °C) prior to use.
 - Plasma or serum specimens containing a precipitate may yield inconsistent test results. Such specimens must be clarified by centrifugation prior to assaying.
- Precautions
 - Anticoagulants including heparin, EDTA and sodium citrate do not affect the test result. Use of other anticoagulants has not been validated. Their use may affect the test result.
 - Use new pipette tips for each specimen in order to avoid cross-contamination of other specimens which could cause erroneous results.

Test procedure (Refer to figure)

- Bring all kit components and specimens to reach a temperature between 15 °C and 30 °C prior to testing.
- Remove the test device from foil pouch and place it on a flat, dry surface. Label the test device with a patient identifier.
- [Using a micropipette]
Dispense 10 µl of plasma or serum specimen or dispense 20 µl of whole blood specimen into the specimen well "S".
OR,
[Using a capillary pipette]
Dispense 20 µl of drawn whole blood specimen into the specimen well "S".
Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S".
⚠ Caution: Do not let bottle nozzle touch device in order to avoid cross-contamination. Hold bottle vertically while dispensing. If you do not hold the bottle vertically, it can lead to inaccurate results. Exactly, 4 drops should be added. Adding more than 4 drops may result in reddish color background or an invalid result.
- As the test begins to work, you will see purple color move across the result window in the center of the test device.
- Interpret test results 10 - 20 minutes after adding assay diluent. Do not read after 20 minutes.
⚠ Caution: If the test result is not legible after 10 minutes due to high background color, read again later but within 20 minutes of adding the diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.

Test interpretation (Refer to figure)

- A colored control line will appear at "C" in the result window to show that the test is working properly.
- The "1" and "2" section of the result window indicates the test result.
- Negative result:**
The presence of only the control line (C) within the result window indicates a negative result.
- Positive result:**
⚠ Caution: The presence of any test line, no matter how faint, the result is considered positive.

- HIV-1 positive:**
 - The presence of both test line 1 (1) and the control line (C) indicates a positive result for anti-HIV-1.
 - If the color intensity of the test line 1 is darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive.
- HIV-2 positive:**
 - The presence of both test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-2.
 - If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive.
- Both HIV-1 and HIV-2 Positive:** The presence of the test line 1 (1), the test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.
⚠ Caution : Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homology in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine the virus type or diagnose a co-infection accurately, you must perform a confirmatory test such as Western Blot etc.
- Invalid result:**
Absence of the control line (C) and/or presence of a pink/purple smear in the result window indicates an invalid result. Instructions may not have been followed correctly or the test may have deteriorated. It is recommended that the specimen be retested using a new test device.

Test limitations

- A positive result may indicate infection with HIV 1/2. Immunochromatographic testing alone cannot be used to diagnose AIDS. An AIDS diagnosis can only be made in a clinical setting if an individual meets the case definition for AIDS established by the Centers for Disease Control (CDC). Positive specimens should be confirmed by a supplemental assay such as ELISA or Western Blot test.
- A negative result does not eliminate the possibility of infection with HIV-1 and/or HIV-2. The specimen may contain low levels of antibodies that cannot be detected by Bioline™ HIV 1/2 3.0 kit. If a test result is negative and clinical symptoms persist, additional testing using other clinical methods is recommended.
- Some known HIV-infected persons taking antiretroviral medication have been shown to produce false negative results when tested by rapid diagnostic tests.
- Where clinical presentation or other data would suggest an inconsistent test result then the individual should be tested by nucleic acid testing (NAT) technologies immediately and/or retested for antibodies to HIV after more than 21 days since the original testing.

Internal quality control

The Bioline™ HIV 1/2 3.0 test device has test line 1, test line 2 and the control line C on the surface of the device. The lines are not visible before applying a specimen. The control line is used for procedural control and shows that the diluent has been applied successfully and that the active ingredients of main components on the strip are functional, but is not a guarantee that the specimen has been properly applied and does not represent a positive specimen control.

Performance characteristics

- Sensitivity and Specificity
699 specimens were tested by Bioline™ HIV 1/2 3.0 and a leading commercially available Anti-HIV 1/2 ELISA kit. The result shows that Bioline™ HIV 1/2 3.0 is well correlated to other commercial ELISA kits. Bioline™ HIV 1/2 3.0 demonstrates a sensitivity of 100% (187/187) and a specificity of 99.8% (511/512).
- Precision
 - INTER RUN : the reproducibility was determined by testing 3 different lots of Bioline™ HIV 1/2 3.0 with 7 different specimens containing different concentrations of antibody. The precision was determined to be 100%.
 - INTRA RUN : the reproducibility was determined by repeating test 3 times with 7 different specimens containing different concentrations of antibody. The precision was determined to be 100%.

Reference	Bioline™ HIV 1/2 3.0			Total Results
Method	Result	Positive	Negative	
Commercial ELISA	Positive	187	0	187
	Negative	1	511	512
Sensitivity (95 % CI)	100 % (98.0 - 100 %)			
Specificity (95 % CI)	99.8 % (98.9 - 100 %)			

Bibliography of suggested reading

- Report of the WHO evaluation(Phase 1) of the SD BIOLINE HIV-1/2 3.0 at WHO collaborating center for Transfusion Transmitted Infectious Department of Institute of Tropical Medicine in Antwerp, Belgium. (2002)
- Zoltan Györi, M.D., and Janos Minarovits, M.D. D.Sc. : Evaluation of the SD BIOLINE HIV-1/2 3.0 rapid test for the detection of antibodies to human immunodeficiency virus (HIV) in sera from European individuals (2008)
- Dr. Heinrich Scheiblaue : EVALUATION OF SD BIOLINE HIV-1/2 3.0 Rapid, MANUFACTURED BY STANDARD DIAGNOSTICS, INC., KOREA (2007)
- Adrian Puren, Deputy Director, Virology : LABORATORY EVALUATION OF HIV RAPID ASSAY (2006)
- Nolan Monica : Résultat de l'évaluation du test rapide SD BIOLINE HIV-1/2 3.0 pour le dépistage du VIH (2005)
- Mr Willie Porau : Summary:HIV Test Kit Evaluation/Validity Study 2006. (2006)
- http://www.who.int/diagnostics_laboratory/evaluations/hiv/en/

<p><i>Product Disclaimer:</i> While every precaution has been taken to ensure the diagnostic ability and accuracy of this product, the product is used outside of the control of the manufacturer and distributor and test results may accordingly be affected by environmental factors and/or user error. The subject of the diagnosis should consult a doctor for further confirmation of the test result.</p> <p><i>Warning:</i> The manufacturers and distributors of this product shall not be liable for any direct, indirect, or consequential losses, liability, claims, costs or damages arising from or related to an incorrect positive or negative diagnosis using this product.</p>
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Date issued : 2023.01
03FK22-01-EN-A0

TECHNICAL SUPPORT CONTACT INFORMATION

For further information, please contact your distributor or contact our support specialists:

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GLOSSARY OF SYMBOLS		
	Temperature limitation	REF Catalog Number
	In vitro diagnostic medical device	Contains sufficient for <n> tests
	Use By	Manufacturer
	Keep dry	Biological Risks
	Date of manufacture	Caution
	Do not reuse	
	Consult instructions for use	
	Do not use if package is damaged	
	Keep away from sunlight	

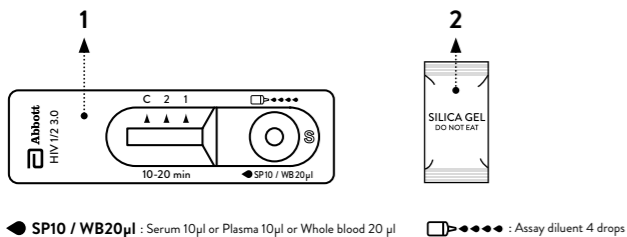
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Bioline™ HIV 1/2 3.0

The 3rd Generation of antibodies to HIV-1/HIV-2 Test

PREPARATION

- Open the package and look for the following:
 - Test device with desiccant in individual foil pouch
 - Assay diluent
 - Instructions for use
 - Capillary pipette (20 µl)
 - Sterile lancet
 - Alcohol swab
- Carefully read the instructions on how to use the Bioline™ HIV 1/2 3.0 test kit.
- Look at the expiration date at the back of the foil pouch. If the expiration date has passed, use another kit. To avoid false results, ensure that the assay diluent used is from the same kit as the new test device.
- Open the foil pouch and look for the following:
 - Test device
 - Desiccant
 Then, label the device with the patient identifier.

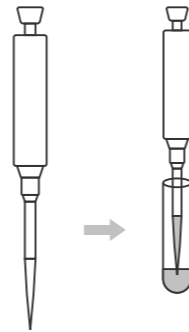


TEST PROCEDURE

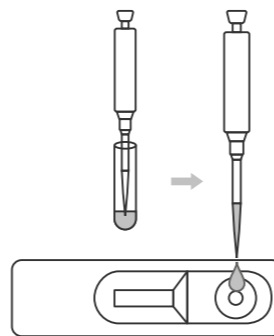
I. Blood (by venipuncture), Plasma or Serum specimen

Specimen collection

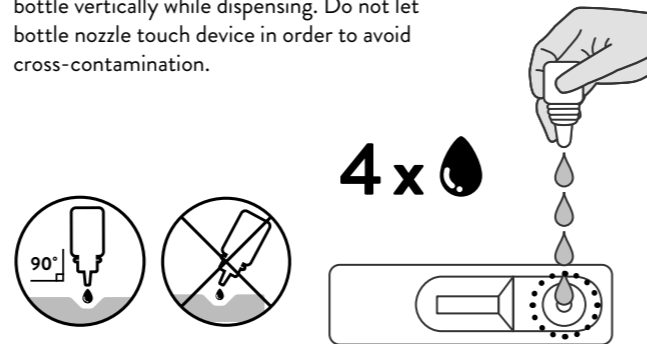
- Using a micropipette draw plasma or serum: 10 µl specimen or whole blood: 20 µl specimen



- Dispense 10 µl of plasma or serum specimen or 20 µl of whole blood specimen into the specimen well marked "S"



- Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.



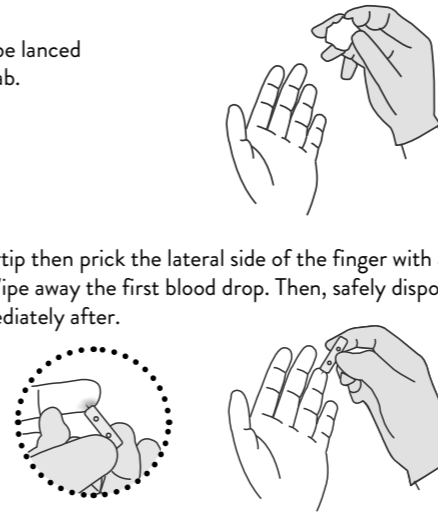
- Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.



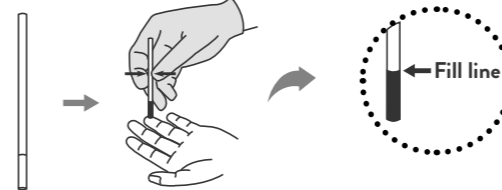
II. Blood specimen (with a lancet)

Specimen collection

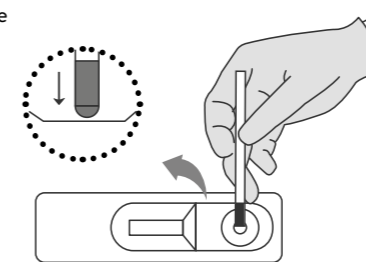
- Clean the area to be lanced with an alcohol swab.
- Squeeze the fingertip then prick the lateral side of the finger with a lancet provided. Wipe away the first blood drop. Then, safely dispose of the lancet immediately after.



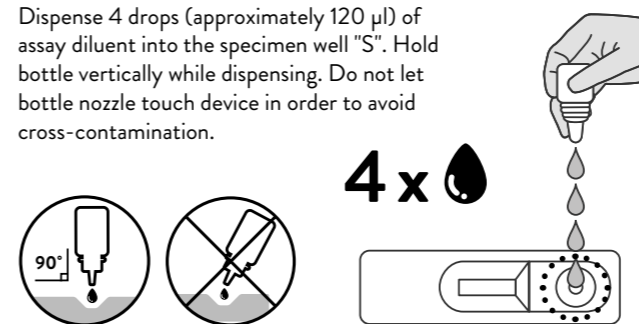
- Immerse the open end of a new capillary pipette (20 µl) in the next blood drop and release the pressure to draw blood into the capillary pipette up to the fill line.



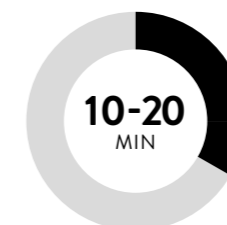
- Dispense 20 µl of drawn whole blood specimen in the specimen well marked "S".



- Dispense 4 drops (approximately 120 µl) of assay diluent into the specimen well "S". Hold bottle vertically while dispensing. Do not let bottle nozzle touch device in order to avoid cross-contamination.



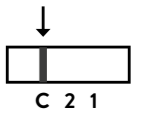
- Interpret the result 10 - 20 minutes after adding assay diluent. Reading outside of this time frame (before 10 min or after 20 min) may provide in false results.



INTERPRETATION

Negative

The presence of only the control line (C) within the result window indicates a negative result.



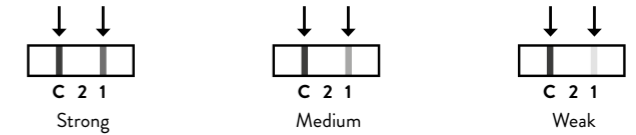
Positive

Caution: The presence of any test line, no matter how faint, the result is considered positive.

HIV-1 Positive

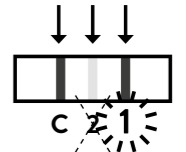
2 LINES

The presence of both test line 1 (1) and the control line (C) indicates a positive result for anti-HIV-1.



3 LINES

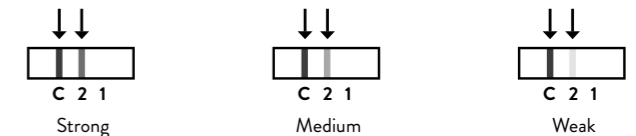
If the color intensity of the test line 1 is darker than one of test line 2 in the result window, you can interpret the result as anti-HIV-1 positive. Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.



HIV-2 Positive

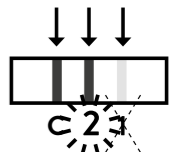
2 LINES

The presence of both test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-2.



3 LINES

If the color intensity of the test line 2 is darker than one of test line 1 in the result window, you can interpret the result as anti-HIV-2 positive. Faint line: By homology in the amino acid sequence of HIV Type-1 and HIV Type-2.



Remark

[Both HIV-1 and HIV-2 Positive]

The presence of the test line 1 (1), the test line 2 (2) and the control line (C) indicates a positive result for anti-HIV-1 and/or anti-HIV-2.

Caution: Although a positive result for anti-HIV-1 and anti-HIV-2 in one patient is a rare case, it's possible as there is an homology in the amino acid sequence between anti-HIV-1 and anti-HIV-2. To determine the virus type or diagnose a co-infection accurately, you must perform a confirmatory test such as Western Blot etc.

Invalid

Absence of the control line (C) and/or presence of a pink/purple smear in the result window indicates an invalid result. Instructions may not have been followed correctly or the test may have deteriorated. It is recommended that the specimen be retested using a new test device.

