

WHO Prequalification of In Vitro Diagnostics PUBLIC REPORT

Product: Bioline HBsAg WB¹
WHO reference number: PQDx 0219-012-00

Bioline HBsAg WB with product code **01FK10W**, manufactured by **Abbott Diagnostics Korea Inc²**, **Rest-of-World (RoW)** regulatory version, was accepted for the WHO list of prequalified in vitro diagnostics and was listed on 22 December 2017.

Summary of WHO prequalification assessment for Bioline HBsAg WB

	Date	Outcome
PQ listing	22-Dec-2017	listed
Dossier review	18-Sep-2017	MR
Site inspection(s) of quality management system	26 – 29-Jun-2017	MR
Product performance evaluation	10-Nov-2016	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 summarized in the following table, and details of each amendment are provided below.

Version	Summary of amendment	Date of report amendment
1.0 to 3.0	Changes to the draft public report before listing.	22-Dec-2016
4.0	Product name was changed from SD BIOLINE HBsAg WB to Bioline HBsAg WB. Manufacturer's name changed from Standard Diagnostics Inc to Abbott Diagnostics Korea Inc.	20-Aug-2020

¹ Product name was changed from SD BIOLINE HBsAg WB to Bioline HBsAg WB.

² Manufacturer's name changed from Standard Diagnostics Inc to Abbott Diagnostics Korea Inc

Intended use:

According to the claim of intended use from Abbott Diagnostics Korea Inc, *“BiolineHBsAg WB is an in vitro immunochromatographic, rapid assay designed for the qualitative detection of Hepatitis B surface antigen, in human serum, plasma (heparin, EDTA and sodium citrate) or venous whole blood (heparin, EDTA and sodium citrate). Bioline HBsAg WB is intended only for professional use as an aid to diagnosis. Reactive specimens should be reflexed for additional testing, either by Enzyme immunoassay (EIA) to identify current HBV infection. This product is intended for use in a population with high HBV prevalence. This test may not be suitable for diagnosis of early infection or blood donation screening. Because false non-reactive results may also arise due to the lack of ability of the assay to detect HBsAg mutants. The performance of Bioline HBsAg WB in infants or children has not been validated.”*

Assay description:

According to the claim of assay description from Abbott Diagnostics Korea Inc, *“the membrane is pre-coated with mouse monoclonal anti-HBsAg pool on the test line region and Mouse monoclonal anti-chicken IgY on the control line region. During testing, the specimen is allowed to react with the colored conjugate (mouse monoclonal anti-HBsAg conjugated gold colloid) which was pre-coated on the test strip. The mixture (mouse monoclonal anti-HBsAg + HBsAg in specimen) then moves upward on the membrane chromatographically by capillary action. For a reactive result, a purple colored line with the antibody-antigen-antibody gold particle complex will form in the test line region of the result window. An absence of this purple-colored line in the test line region suggests a non-reactive result. Regardless of the presence of HBsAg, chicken IgY conjugated gold colloid, pre-coated on the test strip, continues to move across the membrane to immobilized mouse monoclonal anti-chicken IgY, then a purple-colored line at the control line region of the result window appears. The presence of a purple colored line in the control line region serves as 1) verification that sufficient volume of specimen has been added and 2) that proper flow has been obtained.”*

Test kit contents:

Component	30 tests (product code 01FK10W)
Test device: With desiccant in an individual foil pouch.	30
Instructions for use	1

Items required but not provided:

Item
Consumables: Gloves
Durables: N/A
Equipment: Micropipette Timer Biohazard disposal container

Storage:

The test kit should be stored at 1 to 40 °C.

Shelf-life upon manufacture:

24 months.

Warnings/limitations:

See manufacturer's instructions for use.

Prioritization for prequalification

Based on the established eligibility criteria, Bioline HBsAg WB was given priority for WHO prequalification assessment.

Product dossier assessment

Abbott Diagnostics Korea Inc. submitted a product dossier for Bioline HBsAg WB as per the "Instructions for compilation of a product dossier" (PQDx_018 v1). The information (data and documentation) submitted in the product dossier was reviewed by WHO staff and external technical experts (assessors) appointed by WHO.

The manufacturer's responses to the nonconformities found during dossier screening and assessment findings were accepted on 18 September 2017.

Commitments for prequalification:

Commitment for prequalification:

The double reading legend will be removed from the cassette. The change will be implemented in Q2-2018. The updated documentation in relation to this change should be submitted in Q2-2018 or as soon as the change is applied.

WHO will follow-up on implementation of this commitment at the next re-inspection.

Based on the product dossier screening and assessment findings, the product dossier for Bioline HBsAg WB meets WHO prequalification requirements.

Manufacturing site inspection

A comprehensive inspection was performed at the site(s) of manufacture (65, Borahagal-ro, Giheung-gu, Yongin-si, Gyeonggi-do and 46 Hagal-ro 15beon-gil, Giheung-gu, , Yongin-si, Gyeonggi-do) of Bioline HBsAg WB in 26-29 June 2017 as per the “Information for manufacturers on prequalification inspection procedures for the sites of manufacture of diagnostics” (PQDx_014 v1). The inspection found that the manufacturer had an acceptable quality management system and good manufacturing practices in place that would, once the commitments to WHO are fulfilled, ensure the consistent manufacture of a product of good quality.

The manufacturer's responses to the nonconformities found at the time of the inspection were accepted on 19 December 2017.

Commitments for prequalification:

1. Certain process validations to be completed, and applicable production documentation to be updated by June 2018.
2. Stability studies to be performed using samples from routine production batches.
3. Evidence to be provided that the performance of lots submitted for WHO laboratory performance evaluation are equivalent to the performance of routinely manufactured batches by June 2018.

Confirmation of implementation of these commitments will be followed-up at re-inspection.

Based on the site inspection and corrective action plan review, the quality management system for Bioline HBsAg WB meets WHO prequalification requirements, subject to fulfillment of the above commitments.

Manufacturing site inspection

Bioline HBsAg WB is a lateral flow immunochromatographic rapid diagnostic test for the detection of HBsAg in human serum, plasma or whole blood. A volume of 100µL of specimen is needed to perform the assay. This type of assay requires no sophisticated equipment and can therefore be performed in laboratories with limited facilities. Reading of the results can be done visually.

In this limited evaluation on a panel of 514 specimens, we found an initial sensitivity (95% CI of 100% (98.1% – 100%) and an initial specificity (95% CI) of 98.7% (96.8% – 99.7%) compared to the reference assays. The final sensitivity (95% CI) was 100% (98.1% – 100%) and the final specificity (95% CI) was 99.0% (97.2% – 99.8%) compared to the reference assays. The lot to lot variability was acceptable.

Performance characteristics in comparison with an agreed reference standard		
	Initial (95% CI)	Final (95% CI)
Sensitivity %	100% (98.1 – 100%)	100% (98.1 – 100%)
Specificity %	98.7% (96.8 – 99.7%)	99.0% (97.2 – 99.8%)
Invalid rate %	0.2%	
Inter-reader variability %	0.2%	

Additional performance characteristics	
Sensitivity during seroconversion on 6 seroconversion panels in comparison with a benchmark assay; Monolisa Ag HBs Plus (bio-Rad)	Seroconversion sensitivity index of +2, therefore detection is 2 days later than the benchmark assay
Analytical sensitivity on a HBsAg low titer panel in comparison with an agreed reference standard	0 of 14 specimens were correctly classified.
Analytical sensitivity on WHO International Biological Reference Preparation for HBsAg	2.06 IU/ml
Lot to lot variation on a dilution panel in comparison with an agreed reference standard	Acceptable

Key operational characteristics	
Validated specimen types	Serum, plasma (heparin, EDTA and sodium citrate), venous whole blood
Number of steps	1 with precision required
Time to result	20 minutes
Endpoint stability	0 minutes (must be read immediately after 20 minutes)
Internal QC	Yes, reagent addition only
In-use stability of reagents	Use immediately

Labelling

1. Labels
2. Instructions for use

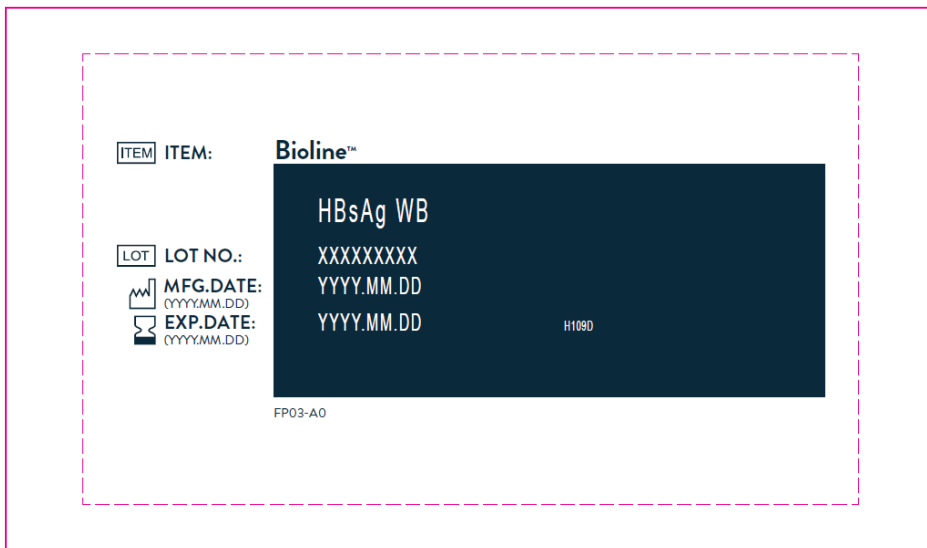
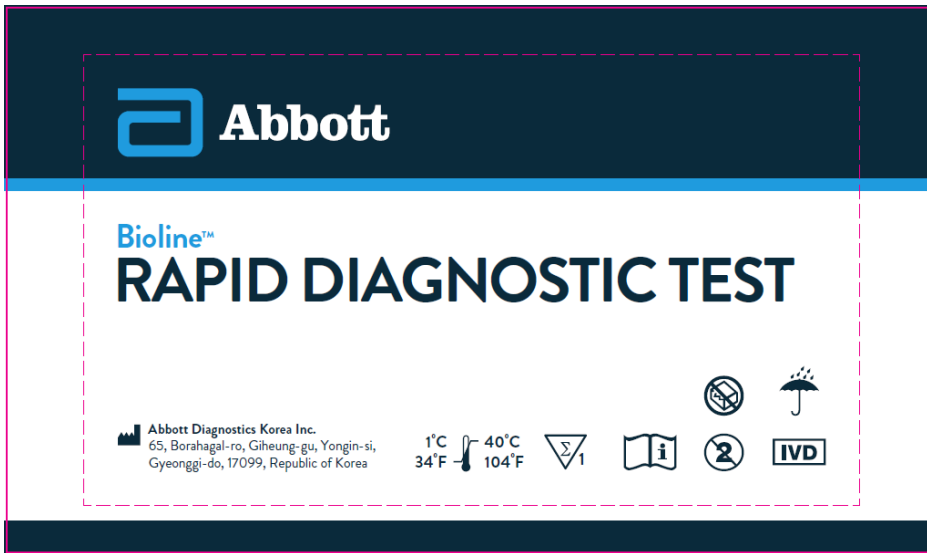
**Note: Labelling has been changed as per site name change and product rebranding. But temporarily labelling of legacy brand (SD BIOLINE) will be used in the market according to registration status in each country.*

1. Labels

1.1 Package box for 01FK10W



1.2 Device pouch for 01FK10W



2. 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.

Bioline[™] HBsAg WB

HBsAg Rapid Test
 Test rapide de détection de l'antigène HBsAg EN UNE ÉTAPE
 Prueba rápida de HBsAg en un paso
 Teste rápido HBsAg num único passo

PREPARATION / PRÉPARATION / PREPARACIÓN / PREPARAÇÃO

- 1** **EN** Open the package and look for the following:
1. Test device with desiccant in individual foil pouch
 2. Instructions for use
- FR** Ouvrir l'emballage et identifier les éléments suivants :
1. Dispositif de test avec agent déshydratant conditionné dans un emballage en aluminium individuel
 2. Mode d'emploi
- ES** Abra el paquete y busque los siguientes elementos:
1. Dispositivo de prueba con desecante en bolsa de papel aluminio individual
 2. Instrucciones de uso
- PT** Abra a embalagem e procure o seguinte:
1. Dispositivo de teste com dessecante em bolsa de folha de alumínio individual
 2. Instruções de utilização

2 **EN** Carefully read the instructions for using the Bioline[™] HBsAg WB test.

FR Lire attentivement le mode d'emploi du test Bioline[™] HBsAg WB.

ES Lea con atención las instrucciones de uso de la prueba Bioline[™] HBsAg WB.

PT Leia cuidadosamente as instruções para utilizar o teste Bioline[™] HBsAg WB.

3 **EN** Look at the expiration date on the back of the foil pouch. If the expiration date has passed, use another kit.

FR Vérifier la date de péremption à l'arrière de l'emballage en aluminium. Si elle est dépassée, utiliser un autre kit.

ES Lea la fecha de vencimiento indicada en la parte posterior de la bolsa. Si la fecha ya ha pasado, use otro kit.

PT Verifique o prazo de validade na parte posterior da bolsa de folha de alumínio. Se o prazo de validade tiver sido ultrapassado, utilize outro kit.

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 Ouvrir l'emballage en aluminium et identifier les éléments suivants :

1. Dispositif de test
2. Agent déshydratant

Apposer ensuite une étiquette indiquant l'identifiant du patient sur le dispositif de test.

ES Abra la bolsa de papel aluminio y busque los siguientes elementos:

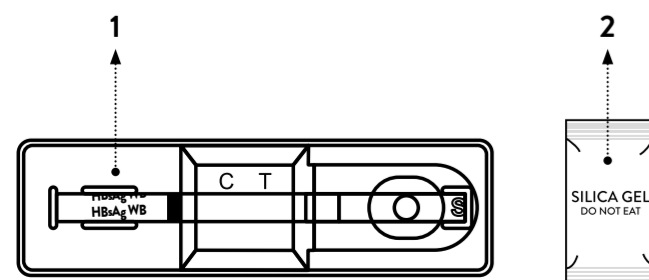
1. Dispositivo de prueba
2. Desecante

Luego, etiquete el dispositivo de prueba con un identificador del paciente.

PT Abra a bolsa de folha de alumínio e procure o seguinte:

1. Dispositivo de teste
2. Dessecante

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



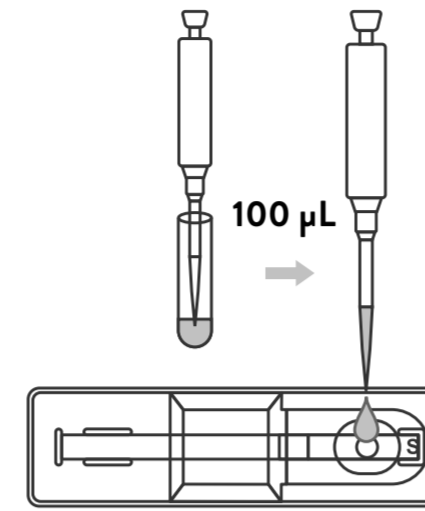
Specimen collection / Prélèvement de l'échantillon / Obtención de la muestra / Colheita de amostras

1 **EN** Take 100 µl of serum, plasma or whole blood specimen using a micropipette. Dispense 100 µl of serum, plasma or whole blood specimen into the specimen well "S".

FR Prélever 100 µl d'échantillon de sérum, de plasma ou de sang total à l'aide d'une micropipette. Déposer 100 µl d'échantillon de sérum, de plasma ou de sang total dans le puits d'échantillon « S ».

ES Con una micropipeta, tome 100 µl de la muestra de suero, plasma o sangre. Instile 100 µl de la muestra de suero, plasma o sangre en el espacio para muestras "S".

PT Tire 100 µL de amostra de soro, plasma ou de sangue total com uma micropipeta. Deite 100 µL de amostra de soro, plasma ou sangue total no poço da amostra "S".

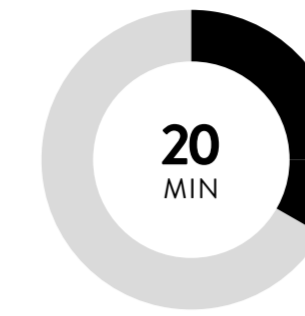


2 **EN** Interpret test results at 20 minutes. Do not read test results after 20 minutes; late readings can yield false results.

FR Interpréter les résultats du test au bout de 20 minutes. Ne pas lire les résultats du test au-delà de 20 minutes, car il est alors possible qu'ils soient erronés.

ES Una vez transcurridos 20 minutos, interprete los resultados. No lea los resultados después de 20 minutos; una lectura tardía puede arrojar resultados falsos.

PT Interprete os resultados do teste após 20 minutos. Não leia os resultados do teste após 20 minutos; as leituras tardias podem gerar resultados falsos.



INTERPRETATION / INTERPRÉTATION / INTERPRETACIÓN / INTERPRETAÇÃO

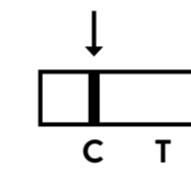
NON-REACTIVE / NON RÉACTIF / NO REACTIVO / NÃO REATIVO

EN The presence of only the control line (C) within the result window indicates a non-reactive result.

FR La présence de la ligne de contrôle uniquement (C) dans la fenêtre de résultat indique un résultat non réactif.

ES Si solo aparece la línea de control (C) en la ventana de resultados, el resultado es no reactivo.

PT A presença apenas da linha de controlo (C) dentro da janela de resultados indica um resultado não reativo.



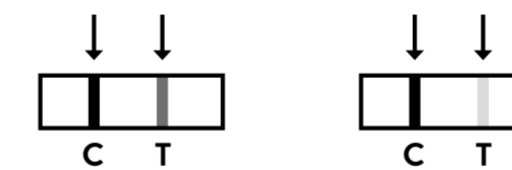
REACTIVE / RÉACTIF / REACTIVO / REATIVO

EN The presence of the test line (T) and the control line (C) within the result window, regardless of which line appears first, indicates a reactive result. **⚠ Caution:** The presence of any test line, no matter how faint, the result is considered reactive.

FR La présence de la ligne de test (T) et de la ligne de contrôle (C) dans la fenêtre de résultat, quelle que soit la ligne apparue en premier, indique un résultat réactif. **⚠ Attention :** si la ligne de test est présente, même très pâle, le résultat est considéré comme réactif.

ES Si aparecen la línea de prueba (T) y la línea de control (C) en la ventana de resultados, independientemente del orden de aparición, el resultado es reactivo. **⚠ Precaución:** La presencia de cualquier línea de prueba, aunque sea de un color débil, indica que el resultado es reactivo.

PT A presença da linha de teste (T) e da linha de controlo (C) dentro da janela de resultados, independentemente da linha que aparecer primeiro, indica um resultado reativo. **⚠ Atenção:** a presença de qualquer linha de teste, mesmo sendo muito tênue, significa que o resultado é considerado reativo.



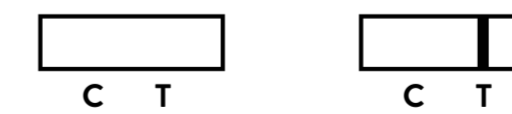
INVALID / NON VALIDE / NO VÁLIDO / INVÁLIDA

EN If the control line (C) is not visible within the result window after performing the test, the result is considered invalid. Instructions may not have been followed correctly or the test kit may have deteriorated. It is recommended that the specimen be retested using a new test device.

FR Si la ligne de contrôle (C) n'est pas visible dans la fenêtre de résultat après la réalisation du test, le résultat est considéré comme non valide. Il se peut que les instructions n'aient pas été suivies correctement ou que le kit de test se soit détérioré. Il est recommandé d'analyser à nouveau l'échantillon à l'aide d'un nouveau dispositif de test.

ES Si no se ve la línea de control (C) en la ventana de resultados después de ejecutar la prueba, se considera que no hay un resultado válido. Esta situación puede deberse a que no se siguieron correctamente las instrucciones o a que el kit de prueba se haya deteriorado. Se recomienda volver a analizar la muestra con un dispositivo de prueba nuevo.

PT Se a linha de controlo (C) não estiver visível dentro da janela de resultados após a realização do teste, o resultado é considerado inválido. As instruções podem não ter sido seguidas corretamente ou o kit de teste pode ter-se deteriorado. Recomenda-se que a amostra seja novamente testada utilizando um novo dispositivo de teste.



Glossary of symbols / Glossaire des symboles / Glosario de símbolos / Glossário de símbolos

1°C / 34°F → 40°C / 104°F	Store at 1 - 40 °C (34 °F - 104 °F) Conserver entre 1 et 40 °C (34 et 104 °F) Almacenar entre 1 y 40 °C (34 °F - 104 °F) Armazenar entre 1 - 40 °C (34 °F - 104 °F)	LOT Lot Number No. de lot Número de Lote Número de lote	Manufacturer Fabricant Fabricante Fabricante
IVD	For in vitro diagnostic use only Pour diagnostic in vitro uniquement Solo para uso de diagnóstico in vitro Somente para uso de diagnóstico in vitro	REF Catalog Number Code produit Número de Referencia Número de Catálogo	Date of manufacture Date de fabrication Fecha de fabricación Data de fabricacao
	Do not reuse Usage unique No Reutilizar Não reutilizar	Instructions for use Attention, voir mode d'emploi Atención, ver Instrucciones de uso Atenção, ver Instruções de uso	Caution Mise en garde Precaución Atenção
	Use By Date de péremption Fecha de caducidad Utilizar até	Keep away from sunlight Conserver à l'abri de la lumière du soleil Manténgase fuera de la luz del sol Manter afastado da luz solar	
	Contains sufficient for <no> tests Permet de réaliser <no> tests Contenido suficiente para <no> pruebas Contém o suficiente para <no> testes	Keep dry Conserver au sec Manténgase seco Conservar seco	
	Do not use if package is damaged Ne pas utiliser si l'emballage est endommagé No utilizar si el envase está dañado Não utilizar se a embalagem estiver danificada	Biological Risks Risques biologiques Riesgos biológicos Riscos biológicos	