

CERTIFICADO DE EXAMEN CE DE DISEÑO
de acuerdo con el Anexo IV, punto 4, de la Directiva 98/79/CE
EC DESIGN-EXAMINATION CERTIFICATE
in accordance with Annex IV, Section 4, Directive 98/79/CE
PRORROGA/EXTENSION — Fecha inicial/ Initial date: 04/12/2008
Fecha de última prórroga/ Last extension date: 27/11/2013

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2008 12 0588 ED	Desde/From 19/11/2018 Hasta/To 18/11/2023	0318

A favor de /In favour of:

Fabricante/Manufacturer:
Nombre/Name: DIA, Pro Diagnostic Bioprobes S.r.l.
Dirección/Address: Via G. Carducci, 27 - 20099- Sesto San Giovanni - Milano (Italy).
Representante autorizado ante la UE/Authorized EU representative:
Nombre/Name: Idem **Dirección/Address:** Idem

Para el producto/For the product:

Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / In Vitro Diagnostic Medical Devices
Grupo genérico/Generic group: Diagnóstico de enfermedades infecciosas / Diagnostic of infectious diseases
Tipo/Type: Especificados en Anexos de este Certificado/Specified in Annexes to this Certificate.

Elaborado en/In the facilities:

DIA, Pro Diagnostic Bioprobes S.r.l.
Via G. Carducci, 27-20099- Sesto San Giovanni - Milano (Italy).

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total N° 2003 12 0388 CT/ This certificate must be accompanied by the EC Full Quality Assurance System Certificate N° 2003 12 0388 CT.

Este certificado es consecuencia de la evaluación de la documentación técnica del diseño contenida en el expediente N° 2003 05 0240, y garantiza que el diseño de los productos descritos cumple los requisitos de la Directiva/ This certificate is issued on the assessment of the design documentation contained in dossier N° 2003 05 0240, and guarantees that the design of the described products fulfil the requirements of the Directive.

Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

Navarro  **agencia española de medicamentos y productos sanitarios**

Fdo. M^a Jesús Lamas Díaz

Firmado digitalmente por: Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018
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Página 1 de 2
CORREO ELECTRÓNICO
on0318@aesmp.es
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ORGANISMO NOTIFICADO 0318

CERTIFICADO DE EXAMEN CE DE DISEÑO
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EC DESIGN-EXAMINATION CERTIFICATE
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A favor de /In favour of:

Fabricante/Manufacturer:
Nombre/Name: DIA, Pro Diagnostic Bioprobes S.r.l.
Dirección/Address: Via G. Carducci, 27 - 20099- Sesto San Giovanni - Milano (Italy).
Representante autorizado ante la UE/Authorized EU representative:
Nombre/Name: Idem **Dirección/Address:** Idem

Tipo de producto / Device type: Reactivos y productos reactivos, calibradores y materiales de control para el diagnóstico de enfermedades infecciosas / Reagents, and reagent products, calibrators and control materials for diagnostic of human infectious diseases.

Clasificación/Classification: Lista A, Anexo II / List A, Annex II

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis B, mediante técnicas de Inmunoabsorción enzimática (ELISA) / Reagents and reagent products for the determination, confirmation and quantification in human specimens of markers of Hepatitis B infection, by Enzyme-linked immunosorbent assay (ELISA)

[NANDO: IVD 0203]

HBs Ag one Version ULTRA ELISA cualitativo / ELISA qualitative

- SAGIULTRA.CE (192 tests)
- SAGIULTRA.CE.96 (96 tests)
- SAGIULTRA.CE.480 (480 tests)
- SAGIULTRA.CE.960 (960 tests)
- SAGIULTRA.CE.DB (192 tests - for Dia.Blood application)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración de conformidad. / This certificate covers all trademarks of these products included by the manufacturer in his declaration of conformity.

Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

Navarro  **agencia española de medicamentos y productos sanitarios**

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PRÓRROGA/EXTENSION — Fecha inicial/ Initial date: 11/12/2003

Fecha de última prórroga/ Last extension date: 27/11/2013

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0390 ED	Desde/From 19/11/2018 Hasta/To 18/11/2023	0318

A favor de/In favour of:

Fabricante/Manufacturer:

Nombre/Name: Dia. Pro Diagnostic Bioprobes S.r.l.

Dirección/Address: Via G. Carducci, 27 - 20099 - Sesto San Giovanni - Milano (Italy).

Representante autorizado ante la UE/Authorized EU representative:

Nombre/Name: Idem **Dirección/Address:** Idem

Tipo de producto / Device type: Reactivos y productos reactivos, calibradores y materiales de control para el diagnóstico de enfermedades infecciosas / *Reagents and reagent products, calibrators and control materials for diagnostic of human infectious diseases.*

Clasificación/Classification: Lista A, Anexo II / *List A, Annex II*

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis B, mediante técnicas de Inmunoabsorción enzimática (ELISA) *Reagents and reagent products for the determination, confirmation and quantification in human specimens of markers of Hepatitis B infection, by Enzyme-linked immunosorbent assay (ELISA)*

(NANDO: TVD 0203)

HBs Ab ELISA cualitativo-cuantitativo / *ELISA qualitative-quantitative*

SAB-CE (96 tests)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración de conformidad. / *This certificate covers all trademarks of these products included by the manufacturer in his declaration of conformity.*

Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

Novanor  **agencia española de medicamentos y productos sanitarios**

Fdo. M^a Jesús Lamas Díaz

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Fecha de la firma: 19/11/2018
Localizador: 62Y6ZAGS9D

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CORREO ELECTRÓNICO

on0318@ampas.es

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ORGANISMO NOTIFICADO 0318

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CERTIFICADO DE EXAMEN CE DE DISEÑO
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A favor de/In favour of:

Fabricante/Manufacturer:

Nombre/Name: DIA. Pro Diagnostic Bioprobes S.r.l.

Dirección/Address: Via G. Carducci, 27 - 20099 - Sesto San Giovanni - Milano (Italy).

Representante autorizado ante la UE/Authorized EU representative:

Nombre/Name: Idem **Dirección/Address:** Idem

Para el producto/For the product:

Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / *In Vitro Diagnostic Medical Devices*

Grupo genérico/Generic group: Diagnóstico de enfermedades infecciosas / *Diagnostic of infectious diseases*

Tipo/Type: Especificados en Anexos de este Certificado/Specified in Annexes to this Certificate.

Elaborado en/In the facilities:

Dia. Pro Diagnostic Bioprobes S.r.l.

Via G. Carducci, 27 - 20099 - Sesto San Giovanni - Milano (Italy).

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total N° 2003 12 0388 CT/ *This certificate must be accompanied by the EC Full Quality Assurance System Certificate N° 2003 12 0388 CT.*

Este certificado es consecuencia de la evaluación de la documentación técnica del diseño contenida en el expediente N° 2003 05 0240, y garantiza que el diseño de los productos descritos cumple los requisitos de la Directiva/ *This certificate is issued on the assessment of the design documentation contained in dossier N° 2003 05 0240, and guarantees that the design of the described products fulfil the requirements of the Directive.*

Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

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MINISTERIO
DE SANIDAD CONSUMO
Y BIENESTAR SOCIAL



CERTIFICADO DE EXAMEN CE DE DISEÑO

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Fecha de última prórroga/ Last extension date: 27/11/2013

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0391 ED	Desde/From 26/11/2018 Hasta/To 18/11/2023	0318

A favor de /In favour of:

Fabricante/Manufacturer:

Nombre/Name: DIA, Pro Diagnostic Bioprobes S.r.l.

Dirección/Address: Via G. Carducci, 27-20099- Sesto San Giovanni – Milano (Italy).

Representante autorizado ante la UE/Authorized EU representative:

Nombre/Name: Idem Dirección/Address: Idem

Para el producto/For the product:

Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / In Vitro Diagnostic Medical Devices

Grupo genérico/Generic group: Diagnóstico de enfermedades infecciosas / Diagnostic of infectious diseases

Tipo/Type: Especificados en Anexos de este Certificado/Specified in Annexes to this Certificate.

Elaborado en/In the facilities:

DIA, Pro Diagnostic Bioprobes S.r.l.

Via G. Carducci, 27-20099- Sesto San Giovanni – Milano (Italy).

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Madrid, 23 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



Fdo. M^a Jesús Lamas Díaz

Firmado originalmente por: Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 23/11/2018

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ORGANISMO NOTIFICADO 0318

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MINISTERIO
DE SANIDAD CONSUMO
Y BIENESTAR SOCIAL



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A favor de /In favour of:

Fabricante/Manufacturer:

Nombre/Name: DIA, Pro Diagnostic Bioprobes S.r.l.

Dirección/Address: Via G. Carducci, 27-20099- Sesto San Giovanni – Milano (Italy).

Representante autorizado ante la UE/Authorized EU representative:

Nombre/Name: Idem Dirección/Address: Idem

Tipo de producto / Device type: Reactivos y productos reactivos, calibradores y materiales de control para el diagnóstico de enfermedades infecciosas / Reagents, and reagent products, calibrators and control materials for diagnostic of human infectious diseases.

Clasificación/Classification: Lista A, Anexo II / List A, Annex II

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis B, mediante técnicas de Inmunoabsorción enzimática (ELISA) / Reagents and reactive products for the determination, confirmation and quantification in human specimens of markers of Hepatitis B infection, by Enzyme-linked immunosorbent assay (ELISA) [NANDO; IVD 0203]

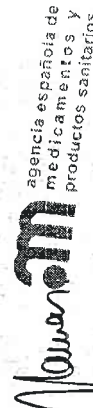
HBc Ab ELISA cualitativo / ELISA qualitative

BCAB CE (96 tests)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración de conformidad. / This certificate covers all trademarks of these products included by the manufacturer in his declaration of conformity.

Madrid, 23 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



Fdo. M^a Jesús Lamas Díaz

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Fecha de la firma: 23/11/2018

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Página 2 de 2

ORGANISMO NOTIFICADO 0318

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Ministerio de Sanidad Consumo y Bienestar Social

CERTIFICADO DE EXAMEN CE DE DISEÑO
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in accordance with Annex IV, Section 4, Directive 98/79/CE
PRÓRROGA/EXTENSION — Fecha inicial/ Initial date: 11/12/2003
Fecha de última prórroga/ Last extension date: 27/11/2013

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0393 ED	Desde/From 19/11/2018 Hasta/To 18/11/2023	0318

A favor de /In favour of:

Fabricante/Manufacturer:
Nombre/Name: DIA. Pro Diagnostic Bioprobes S.r.l.
Dirección/Address: Via G. Carducci, 27 - 20099- Sesto San Giovanni - Milano (Italy).
Representante autorizado ante la UE/Authorized EU representative:
Nombre/Name: Idem Dirección/Address: Idem

Para el producto/For the product:
Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / In Vitro Diagnostic Medical Devices
Grupo genérico/Generic group: Diagnóstico de enfermedades infecciosas / Diagnostic of infectious diseases
Tipo/Type: Especificados en Anexos de este Certificado/Specified in Annexes to this Certificate.

Elaborado en/In the facilities:
DIA. Pro Diagnostic Bioprobes S.r.l.
Via G. Carducci, 27 - 20099- Sesto San Giovanni - Milano - (Italy).

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total N° 2003 12 0388 CT/ This certificate must be accompanied by the EC Full Quality Assurance System Certificate N° 2003 12 0388 CT.

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Madrid, 19 de noviembre de 2018
DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



Fdo. M^a Jesús Lamas Díaz

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Página 1 de 2
ORGANISMO NOTIFICADO 0318
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Fax: (+34) 91 822 52 86



Ministerio de Sanidad Consumo y Bienestar Social

CERTIFICADO DE EXAMEN CE DE DISEÑO
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Fabricante/Manufacturer:
Nombre/Name: Dia. Pro Diagnostic Bioprobes S.r.l.
Dirección/Address: Via G. Carducci, 27 - 20099- Sesto San Giovanni - Milano (Italy).
Representante autorizado ante la UE/Authorized EU representative:
Nombre/Name: Idem Dirección/Address: Idem

Tipo de producto / Device type: Reactivos y productos reactivos, calibradores y materiales de control para el diagnóstico de enfermedades infecciosas / Reagents, and reagent products, calibrators and control materials for diagnostic of human infectious diseases.

Clasificación/Classification: Lista A, Anexo II / List A, Annex II

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis D, mediante técnicas de Inmunoabsorción enzimática (ELISA)/ Reagents and reactive products for the determination, confirmation and quantification in human specimens of markers of Hepatitis D infection, by Enzyme-linked immunosorbent assay (ELISA)

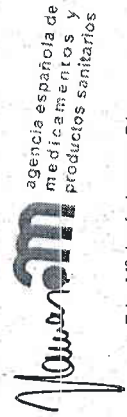
[NANDO: IVD 0203]

HDV Ab ELISA cualitativo / ELISA qualitative

- DAB-CE (96 tests)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración de conformidad. / This certificate covers all trademarks of these products included by the manufacturer in his declaration of conformity.

Madrid, 19 de noviembre de 2018
DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



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Página 2 de 2
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Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0392 ED	Desde/From 19/11/2018 Hasta/To 18/11/2023	0318

A favor de /In favour of:

Fabricante/Manufacturer:
Nombre/Name: DIA, Pro Diagnostic Bioprobes S.r.l.
Dirección/Address: Via G. Carducci, 27 - 20099- Sesto San Giovanni – Milano (Italy).
Representante autorizado ante la UE/Authorized EU representative:
Nombre/Name: Idem Dirección/Address: Idem

Para el producto/For the product:

Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / *In Vitro Diagnostic Medical Devices*
Grupo genérico/Generic group: Diagnóstico de enfermedades infecciosas / *Diagnostic of infectious diseases*
Tipo/Type: Especificados en Anexos de este Certificado/Specified in Annexes to this Certificate.

Elaborado en/In the facilities:

DIA, Pro Diagnostic Bioprobes S.r.l.
Via G. Carducci, 27 - 20099- Sesto San Giovanni – Milano (Italy).

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Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS


agencia española de
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Nombre/Name: Idem Dirección/Address: Idem

Tipo de producto / Device type: Reactivos y productos reactivos, calibradores y materiales de control para el diagnóstico de enfermedades infecciosas / *Reagents, and reagent products, calibrators and control materials for diagnostic of human infectious diseases.*

Clasificación/Classification: Lista A, Anexo II / *List A, Annex II*

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis C, mediante técnicas de Inmunoabsorción enzimática (ELISA)/ Reagents and reagent products for the determination, confirmation and quantification in human specimens of markers of Hepatitis C infection, by Enzyme-linked immunosorbent assay (ELISA)

[NANDO: IVD 0203]

HCV Ab ELISA cualitativo / *ELISA qualitative*

- CVAB.CE (192 tests)
- CVAB.CE.96 (96 tests)
- CVAB.CE.480 (480 tests)
- CVAB.CE.960 (960 tests)
- CVAB.CE.DB (192 tests - for Dia.Blood application)

Este certificado ampara todas las marcas de estos productos incluidas por el fabricante en su declaración de conformidad. / *This certificate covers all trademarks of these products included by the manufacturer in his declaration of conformity.*

Madrid, 19 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS


agencia española de
medicamentos y
productos sanitarios

Fdo. M^a Jesús Lamas Díaz



Dia.Pro
Diagnostic
BioProbes

EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	HAV IgM CODE: AVM.CE (96 tests)
CLASSIFICATION	GENERAL IVD
CONFORMITY ASSESSMENT ROUTE	SELF CERTIFICATION

WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS
THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC
FOR IN VITRO DIAGNOSTIC DEVICES.

ISO CERTIFICATE	UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY AEMPS (AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS)
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PLACE & DATE OF FIRST ISSUE	MILANO – SEPTEMBER 2003
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – MARCH 2019
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018



Dia.Pro
Diagnostic
BioProbes

EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	HP IgG CODE: HPG.CE (96 tests)
CLASSIFICATION	GENERAL IVD
CONFORMITY ASSESSMENT ROUTE	SELF CERTIFICATION

WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS
THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC
FOR IN VITRO DIAGNOSTIC DEVICES.

ISO CERTIFICATE	UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY AEMPS (AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS)
-----------------	---

PLACE & DATE OF FIRST ISSUE	MILANO – MARCH 2004
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – MARCH 2019
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018

Сертификат

mdc medical device certification GmbH

удостоверяет, что на предприятии

ВЕКТОР



АО «Вектор-Бест»

**630559, Новосибирская область, р.п. Кольцово,
Научно-производственная зона, корпус 36, к. 211,
Российская Федерация**

с производственными площадками согласно приложению к Сертификату

применительно к областям

**проектирование и разработка, производство и реализация
медицинских изделий in-vitro диагностики
(ПЦР, ИФА, биохимия)**

была введена и применяется

СИСТЕМА УПРАВЛЕНИЯ КАЧЕСТВОМ

Проведенная проверка системы управления качеством показала,
что данная система соответствует требованиям стандарта:

EN ISO 13485

**Изделия медицинские – Системы менеджмента качества –
Регулирующие системные требования**

EN ISO 13485:2016 + AC:2016 - ISO 13485:2016

Дата выдачи	2018-07-13
Срок действия до	2020-07-03
Регистрационный №	D1213100017
Отчет №	P18-00489-117996
Штутгарт, Германия	2018-07-13



Руководитель сертификационного органа



mdc medical device certification GmbH
Kriegerstraße 6
D-70191 Stuttgart, Germany
Phone: +49-(0)711-253597-0
Fax: +49-(0)711-253597-10
Internet: <http://www.mdc-ce.de>

For electronic publication only

Приложение к Сертификату

№ D1213100017

от 2018-07-13

Стр. 1 из 1

Месторасположение	Область действия
АО «Вектор-Бест», ул. Арбузова, 1/1, 630117, г. Новосибирск, Российская Федерация	проектирование и разработка, производство и реализация медицинских изделий in vitro диагностики
АО «Вектор-Бест», 630559, Новосибирская область, р.п. Кольцово, Научно-производственная зона, корпус 36, Российская Федерация	проектирование и разработка, производство медицинских изделий in vitro диагностики
АО «Вектор-Бест», ул. Пасечная, 3, 630117, г. Новосибирск, Российская Федерация	проектирование и разработка, производство медицинских изделий in vitro диагностики



Руководитель сертификационного органа



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For electronic publication only

EC DECLARATION OF CONFORMITY

ZAO "Vector-Best" hereby ensures under own responsibility and declares that the products listed on pages 2-4 are in conformity with applicable provisions and fulfill the essential requirements of Annex I Directive 98/79/EC of 27 October 1998 regarding in vitro diagnostic medical devices.

Classification of products:

Other devices (all devices except Annex II and self-testing devices)

Conformity assessment procedure:

Annex III (not including section 6).

Manufacturer:

ZAO "Vector-Best"
Address: AHC, Koltsovo,
Novosibirsk Region, 630559, Russia,
Tel.: +7 (383) 363 20 60,
Fax: +7 (383) 363 35 55

European authorized representative:

Bioron GmbH,
Rheinhorststr. 18, D-67071
Ludwigshafen, Germany,
tel.: +49 (0) 621 5720 915,
fax: +49 (0) 621 5720 916

Date: 2013/04/12

Murat Khusainov
General Director ZAO «Vector-Best»



No.	Product name	Identification data	REF
1.	Vectohep A-IgM	ELISA kit for determination of IgM to hepatitis A virus	D-0352
2.	Vectohep A-IgG	ELISA kit for quantitative and qualitative determination of IgG to hepatitis A virus	D-0362
3.	Vectohep TTV-IgG	ELISA kit for determination of IgG to TT virus	D-0802
4.	Vectohep E-IgG	ELISA kit for determination of IgG to hepatitis E virus	D-1056
5.	Vectohep E-IgM	ELISA kit for determination of IgM to hepatitis E virus	D-1058
6.	Vectohep G-IgG	ELISA kit for determination of IgG to hepatitis G virus	D-1252
7.	LymeBest-IgG	ELISA kit for determination of IgG to infectious borreliosis agents	D-1452
8.	LymeBest-IgM	ELISA kit for determination of IgM to infectious borreliosis agents	D-1454
9.	RecombiBest antipallidum-IgG	ELISA kit for determination of IgG to Treponema pallidum	D-1852
10.	RecombiBest antipallidum-total antibodies	ELISA kit for determination of total antibodies to Treponema pallidum	D-1856
11.	RecombiBest antipallidum-IgM	ELISA kit for determination of IgM to Treponema pallidum	D-1858
12.	RecombiBest antipallidum-total antibodies	ELISA kit for determination of total antibodies to Treponema pallidum	D-1857
13.	VectoHSV-1,2 - IgG	ELISA kit for determination of IgG to herpes simplex virus types 1 and 2	D-2152
14.	VectoHSV - IgM	ELISA kit for determination of IgM to herpes simplex virus types 1 and 2	D-2154
15.	VectoHHV-8 - IgG	ELISA kit for determination of IgG to human herpes virus type 8	D-2160
16.	VectoHHV-6 - IgG	ELISA kit for determination of IgG to human herpes virus type 6	D-2166
17.	Ureaplasma urealyticum - IgG-EIA-BEST	ELISA kit for determination of IgG to Ureaplasma urealyticum antigens	D-2254
18.	Ureaplasma urealyticum - IgA-EIA-BEST	ELISA kit for determination of IgA to Ureaplasma urealyticum antigens	D-2258
19.	VectoParotitis-IgG	ELISA kit for determination of IgG to parotitis virus	D-2602
20.	VectoParotitis-IgM	ELISA kit for determination of IgM to parotitis virus	D-2604
21.	Toxocara-IgG-EIA-BEST	ELISA kit for determination of IgG to toxocara antigens	D-2752
22.	Opisthorchiasis - IgG-EIA-BEST	ELISA kit for determination of IgG to opisthorchiasis antigens	D-2952
23.	Echinococcus-IgG-EIA-BEST	ELISA kit for determination of IgG to Echinococcus	D-3356

	antigens	
24.	Ascarid-IgG-EIA-BEST	D-3452
25.	Lambliia-antibodies-EIA-BEST	D-3552
26.	Lambliia-IgM-EIA-BEST	D-3554
27.	Lambliia-antigen-EIA-BEST	D-3556
28.	Helicobacter pylori-CagA-antigen-EIA-BEST	D-3752
29.	TSH-EIA-BEST	X-3952
30.	T3 total-EIA-BEST	X-3954
31.	T4 total-EIA-BEST	X-3956
32.	Anti-TPO-EIA-BEST	X-3968
33.	PAPP-A-EIA-BEST	D-4160
34.	Mycoplasma hominis-IgG-EIA-BEST	D-4352
35.	Mycoplasma hominis-IgA-EIA-BEST	D-4358
36.	Mycoplasma pneumoniae-IgG-EIA-BEST	D-4362
37.	Mycoplasma pneumoniae-IgM-EIA-BEST	D-4366
38.	Vectocrimmean - CHF - IgG	D-5052
39.	Vectocrimmean - CHF - IgM	D-5054
40.	CEA-EIA-BEST	T-8454
41.	AFP-EIA-BEST	T-8456
42.	CA-125-EIA-BEST	T-8466
43.	CA 19-9-EIA-BEST	T-8470
44.	CA 15-3-EIA-BEST	T-8472
45.	NSE-EIA-BEST	T-8476

		ELISA kit for determination of concentration of	
46.	Ferritin-EIA-BEST	ELISA kit for determination of concentration of ferritin	T-8552
47.	IgE total-EIA-BEST	ELISA kit for determination of concentration of total IgE	A-8660
48.	IgG total-EIA-BEST	ELISA kit for determination of concentration of total IgG	A-8662
49.	IgM total-EIA-BEST	ELISA kit for determination of concentration of total IgM	A-8664
50.	IgA total-EIA-BEST	ELISA kit for determination of concentration of total IgA	A-8666
51.	Gamma-Interferon-EIA-BEST	ELISA kit for determination of concentration of gamma-interferon	A-8752
52.	Interleukine-4-EIA-BEST	ELISA kit for determination of concentration of Interleukine-4	A-8754
53.	Alpha-TNF-EIA-BEST	ELISA kit for determination of concentration of alpha-tumor necrosis factor	A-8756
54.	Alpha-Interferon-EIA-BEST	ELISA kit for determination of concentration of alpha-interferon	A-8758
55.	Interleukine-6-EIA-BEST	ELISA kit for determination of concentration of Interleukine-6	A-8768
56.	Interleukine-2-EIA-BEST	ELISA kit for determination of concentration of Interleukine-2	A-8772
57.	Procalcitonin-EIA-BEST	ELISA kit for determination of concentration of procalcitonin	A-9004
58.	NTproBNP-EIA-BEST	ELISA kit for determination of concentration of N-terminal prohormone of brain natriuretic peptide	A-9102
59.	Troponin I-EIA-BEST	ELISA kit for determination of concentration of troponin I	A-9106



DECLARATION OF CONFORMITY

- 1) Manufacturer (Name, department): **Monobind Inc.**
Address: **100 North Pointe, LAKE FOREST, CA 92630, UNITED STATES**
and
- 2) European authorized representative: **CEpartner4U BV**,
Address: **ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS**,
(on product labels printed as: **CEpartner4U**, **ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS**, **www.cepartner4u.eu**)
- 3) Product(s) (name, type or model/batch number, etc.):

Immunoassay products;
ELISA,
CLIA,
Control,
Instruments
(see appendix)

- 4) The product(s) described above is in conformity with:

Title	Document No.
In vitro Diagnostic Medical Devices Directive	98/79/EC

- 5) Additional information (Conformity procedure, Notified Body, CE certificate, Registration nr., etc.):
Conformity assessment procedure for CE marking: **In vitro Diagnostic Medical Device Directive**,
Annex III
Registration nr. : **NL- CA002-22758 and NL- CA002-22762**

Lake Forest, USA; 2013-09-16
(Place & date of issue (yyyy-mm-dd))

Tony Shatola
Tony Shatola; QA Director, Monobind Inc.
(name; function and signature of manufacturer)

Olga Teirlinck
Olga Teirlinck; Consultant, CEpartner4U BV
(name; function and signature of authorized representative)

Appendix

List of devices.

Device types	Item# AccuBind® ELISA Microwalls	Item# AccuBind® CLIA Microwalls	Item# QSure® Control	Item# Instrument	EDMS code	Risk Class	First date of CE-marking
Thyroid							
Total Triiodothyronine (T3) Test System	125-300	175-300			12.04.01.05.00	Low	2005-11-11
Free Triiodothyronine (T3) Test System	1325-300	1375-300			12.04.01.01.00	Low	2005-11-11
Thyroxine (T4) Test System	225-300	275-300			12.04.01.07.00	Low	2005-11-11
Free Thyroxine (T4) Test System	1225-300	1275-300			12.04.01.02.00	Low	2005-11-11
Thyrotropin (TSH) Test System	325-300	375-300			12.04.01.11.00	Low	2005-11-11
Rapid TSH Test System	8025-300	8075-300			12.04.01.11.00	Low	2010-06-29
T3-Uptake (T3U) Test System	525-300	575-300			12.04.01.06.00	Low	2005-11-11
Thyroxine-Binding Globulin (TBG) Test System	3525-300	3575-300			12.04.01.09.00	Low	2005-11-11
Thyroglobulin (Tg) Test System	2225-300	2275-300			12.04.01.08.00	Low	2005-11-11
Total Thyroxine (T4), Total Triiodothyronine (T3) & Thyroid Stimulating Hormone (TSH) Thyroid Panel (VAST) Test System	8025-300	8075-300			12.04.01.01.00	Low	2005-11-11
Total Triiodothyronine (T3 SBS) Test System	8125-300	8175-300			12.04.01.01.00	Low	2010-06-29
Total Thyroxine (T4 SBS) Test System	8225-300	8275-300			12.04.01.01.00	Low	2010-06-29
Free Thyroxine (T4), Free Triiodothyronine (T3) & Thyroid Stimulating Hormone Free Thyroid Panel (VAST) Test System	7025-300	7075-300			12.04.01.01.00	Low	2010-06-29
Neonatal Thyroid & Genetics							
Neonatal TSH (N-TSH) Test System	3425-300	3475-300			12.04.01.90.00	Low	2005-11-11
Neonatal (N-T4) Thyroxine Test System	2625-300	2675-300			12.04.01.12.00	Low	2005-11-11
Neonatal 17OHP (N-17OHP) Test System	5525-300	5575-300			12.05.01.07	Low	2008-02-01
Neonatal TBG (N-TBG) Test System	8925-300	8975-300			12.04.01.09.00	Low	2013-09-16
Autoimmune Thyroid							
Anti-Thyroglobulin (Anti-Tg) Test System	1025-300	1075-300			12.10.03.04.00	Low	2005-11-11
Anti-Thyroperoxidase (Anti-TPO) Test System	1125-300	1175-300			12.10.03.01.00	Low	2005-11-11
Fertility & Prenatal							
Luteinizing Hormone (LH) Test System	625-300	675-300			12.05.01.05.00	Low	2005-11-11
Follicle Stimulating Hormone (FSH) Test System	425-300	475-300			12.05.01.04.00	Low	2005-11-11
Prolactin Hormone (PRL) Test System	725-300	775-300			12.05.01.06.00	Low	2005-11-11
Prolactin Hormone Sequential (PRLs) Test System	6025-300	6075-300			12.05.01.06.00	Low	2005-11-11
B-Human Chorionic Gonadotropin (hCG) Test System	825-300	875-300			12.05.02.05.00	Low	2005-11-11
B-Human Chorionic Gonadotropin Extended Range (Ext. Range hCG) Test System	8825-300	8875-300			12.05.02.05.00	Low	2013-09-16
Rapid B-Human Chorionic Gonadotropin (Rapid	3325-300				12.05.02.05.00	Low	2005-11-11

Device types	Item# AccuBind® ELISA Microtiter	Item# AccuLine® CLIA Microtiter	Item# QSure® Control	Item# Instum. ent.	EDMS code	Risk Class	First date of CE-Marking
hCG Test System							
Human Chorionic Gonadotropin (hCG) : Human Prolactin (hPL), Human Luteinizing Hormone (hLH), Follicle Stimulating Hormone (FSH) Fertility Panel (VAST) Test System	8325-300	8375-300			12.05.01.90.00	Low	2005-08-24
Alpha-Fetoprotein (AFP), Human Chorionic Gonadotropin (hCG), Unconjugated Estiol (u- E3) Triple Screen (VAST) Test System	8525-300	8575-300			12.05.01.90.00	Low	2010-06-29
Pregnancy Associated Plasma Protein - A (PAPP-A) Test System	7925-300	7975-300			12.05.02.10.00	Low	2013-09-16
Steroid							
Cortisol Test System	3625-300	3675-300			12.06.02.04.00	Low	2005-11-11
DHEA-S Test System	5125-300	5175-300			12.05.01.02.00	Low	2010-06-29
Dehydroepiandrosterone (DHEA) Test System	7425-300	7475-300			12.05.01.02.00	Low	2011-09-26
Estradiol (E2) Test System	4925-300	4975-300			12.05.01.03.00	Low	2010-06-29
Unconjugated Estiol (u-E3) Test System	5025-300	5075-300			12.05.02.02.00	Low	2010-06-29
Progesterone Test System	4825-300	4875-300			12.05.01.06.00	Low	2010-06-29
Sex Hormone Binding Globulin (SHBG) Test System	9125-300	9175-300			12.05.01.09.00	Low	2013-09-16
Testosterone Test System	3725-300	3775-300			12.05.01.10.00	Low	2007-11-01
Free Testosterone Test System	5325-300	5375-300			12.05.01.10.00	Low	2010-06-29
17α-OH Progesterone Test System	5225-300	5275-300			12.05.01.07.00	Low	2010-06-29
17α-OH Progesterone - SI Test System	9525-300	9575-300			12.05.01.07.00	Low	2010-10-18
Growth & Bone Metabolism							
Growth Hormone (hGH) Test System	1725-300	1775-300			12.06.04.02.00	Low	2005-11-11
Parathyroid Hormone (PTH) Test System	9225-300	9275-300			12.06.03.13.00	Low	2011-09-26
25-Hydroxyvitamin D3 (Vitamin D3) Test System	7725-300	7775-300			12.06.03.10.00	Low	2011-09-26
Diabetes							
Insulin Test System	2425-300	2475-300			12.06.01.03.00	Low	2005-11-11
Rapid Insulin Test System	5825-300				12.06.01.03.00	Low	2010-06-29
C-Peptide Test System	2725-300	2775-300			12.06.01.01.00	Low	2005-11-11
Insulin - C-Peptide (VAST)	7325-300	7375-300			12.06.01.03.00	Low	2005-11-11
Cardiac Markers							
CK-MB Test System	2925-300	2975-300			12.13.01.02.00	Low	2005-11-11
Troponin I (cTnI) Test System	3825-300	3875-300			12.13.01.07.00	Low	2005-11-11
Digoxin (DIG) Test System	925-300	975-300			12.08.01.01.00	Low	2005-11-11
High Sensitivity CRP (hs-CRP) Test System	3125-300	3175-300			12.13.01.09.00	Low	2005-11-11
Myoglobin Test System	3225-300	3275-300			12.13.01.05.00	Low	2005-11-11

Device types	Item# AccuBind® ELISA Microtiter	Item# AccuLine® CLIA Microtiter	Item# QSure® Control	Item# Instum. ent.	EDMS code	Risk Class	First date of CE-Marking
Infectious Diseases							
Anti-H. Pylori IgG Test System	1425-300	1475-300			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgM Test System	1525-300	1575-300			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgA Test System	1625-300	1675-300			15.01.04.03.00	Low	2005-11-11
Cancer Markers							
Alpha-Fetoprotein (AFP) Test System	1925-300	1975-300			12.03.90.01.00	Low	2005-11-11
CA-125 Test System	3025-300	3075-300			12.03.01.06.00	Low	2005-11-11
CA-15-3 Test System	5625-300	5675-300			12.03.01.02.00	Low	2010-06-29
CA-19-9 Test System	3925-300	3975-300			12.03.01.03.00	Low	2005-11-11
Carcinoembryonic Antigen (CEA) Test System	1825-300	1875-300			12.03.01.31.00	Low	2005-11-11
Next Generation Carcinoembryonic Antigen (CEA) Test System	4625-300	4675-300			12.03.01.31.00	Low	2010-06-29
Free β-Subunit Human Chorionic Gonadotropin (βhCG) Test System	2025-300	2075-300			12.03.01.90.00	Low	2005-11-11
Allergy & Anemia							
Ferritin Test System	2825-300	2875-300			12.07.01.02.00	Low	2005-11-11
Folate Test System	7525-300	7575-300			12.07.01.03.00	Low	2010-06-29
Immunoglobulin E (IgE) Test System	2525-300	2575-300			12.02.01.02.00	Low	2005-11-11
Transferrin Soluble Receptor (sTfR) Test System	8625-300	8675-300			12.07.01.06.00	Low	2010-06-29
Vitamin B-12 (B12) Test System	7625-300	7675-300			12.07.02.04.00	Low	2011-09-26
Folate, Vitamin B-12 (VAST) Test System	7825-300	7875-300			12.07.01.00.00	Low	2013-09-16
Miscellaneous Controls							
Anti-Thyroglobulin (Anti-Tg), Anti- Thyroperoxidase (Anti-TPO) Control - Positive & Negative			ALT-101		12.50.01.16.00	Low	2010-06-29
High Level Fertility Control - Single Level - Progesterone, Estradiol, Human Chorionic Gonadotropin			FC-300		12.50.01.16.00	Low	2010-06-29
Maternal Control - Tri Level - Human Chorionic Gonadotropin, Free Beta Human Chorionic Gonadotropin Subunit, Alpha Feto Protein, Estradiol			MC-300		12.50.01.16.00	Low	2010-06-29
Thyroglobulin (Tg) Control - Tri Level			TG-300		12.50.01.16.00	Low	2010-06-29
H. Pylori IgG Control - Positive & Negative			HPY-IgG-300		12.50.01.16.00	Low	2010-06-29
H. Pylori IgM Control - Positive & Negative			HPY-IgM-300		12.50.01.16.00	Low	2013-09-16
H. Pylori IgA Control - Positive & Negative			HPY-IgA-300		12.50.01.16.00	Low	2013-09-16
Thyroid Binding Globulin (TBG) Control - Tri Level			TBG-300		12.50.01.16.00	Low	2013-09-16
Miscellaneous Instruments							
Autochem ELISA & CLIA Analyzer				IN006	21.02.10.01	Low	2010-06-29
Autochem Generation 2 ELISA & CLIA Analyzer				IN005-2	21.02.10.01	Low	2013-09-16
Lumax CLIA Analyzer				IN001	21.02.10.01	Low	2006-06-24
Nico-Lumax CLIA Analyzer				IN010	21.02.10.01	Low	2011-09-26



Page: 5 of 5

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Page: 5 of 5

Device types	Item# Accutest® ELISA Microwells	Item# Accutest® ELISA Microwells	Item# QSure® Control	Item# Instrum ent	EDMS code	Risk Class	First date of CE-marking
Impulse 2 CLIA Analyzer				IN005	21.02.10.01	Low	2006-08-24
Impulse 3 CLIA Analyzer				IN007	21.02.10.01	Low	2010-06-29
Lumax86 CLIA Analyzer				IN004	21.02.10.01	Low	2007-03-01
Lumatic CLIA Analyzer				IN008	21.02.10.01	Low	2011-09-26
Eldex 3.8 ELISA Analyzer				IN003	21.02.10.01	Low	2007-09-10
Neo-Eldex ELISA Analyzer				IN009	21.02.10.01	Low	2011-09-26
PrisMatic ELISA Analyzer				IN013	21.02.10.01	Low	2013-09-16
Plate Washer Microplate Washer				IN002	21.02.10.01	Low	2010-06-29



EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
ASO Latex kit	031100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.



Eddy Velthuis
Technical Director



File No A12241:
ISO 13465:2003; ISO 9001:2008

Lorne Laboratories Limited	Tel: +44 (0) 118 921 2264
Unit 1 Cutbush Park Industrial Estate	Fax: +44 (0) 118 986 4518
Danehill, Lower Earley	Email: info@lornelabs.com
Berkshire RG6 4UT United Kingdom	www.lornelabs.com

Registered office as above. Registered in England No. 04540797. VAT No. 800 3655 66

EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
RF Latex kit	830100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.



Eddy Velthuis
Technical Director



File No A12241:
ISO 13485:2003; ISO 9001:2008

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Danehill, Lower Earley
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Email: info@lornelabs.com
www.lornelabs.com

Registered office as above. Registered in England No. 01540797. VAT No. 800 3655 66

EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
CRP Latex kit	850100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.



Eddy Velthuis
Technical Director



File No A12241:
ISO 13485:2003; ISO 9001:2008

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Berkshire RG6 4UT United Kingdom | www.lornelabs.com

Registered office as above. Registered in England No. 04540797, VAT No. 890 3655 66



CERTIFICATE OF REGISTRATION

Lorne Laboratories Ltd

Unit 1 Cutbush Park Industrial Estate
Danehill
Lower Earley
Berkshire RG6 4UT UNITED KINGDOM

UL LLC®(UL) issues this certificate to the Firm named above, after assessing the Firm's quality system and finding it in compliance with:

ISO 13485:2016

EN ISO 13485:2016

The manufacture of in vitro diagnostic blood grouping reagents. The purchase for resale of in vitro diagnostic serology test kit.

Authorized by

Michael J. Windler, P.E.

Manager of Global Regulatory Service
Distinguished Member of the Technical Staff
UL Life and Health Sciences
UL LLC



Check Certificate
Status: [here](#)

File Number A12241
Certificate 1458.180626
Initial Issue June 26, 2018

Cycle Start Date June 26, 2018
Effective Date June 26, 2018
Expiry Date May 22, 2020

This quality system registration is included in UL's Directory of Registered Firms and applies to the provision of goods and/or services as specified in the scope of registration from the address(es) shown above. By issuance of this certificate the firm represents that it will maintain its registration in accordance with the applicable requirements. This certificate is not transferable and remains the property of UL LLC.



00-MB-S0043 Issue 15.0

UL LLC
333 Pfingsten Road
Northbrook, IL 60062-2096
USA

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EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product Name	Catalogue Number
Anti-A Monoclonal	600010

has been classified as List A (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC) and the Commission Decision on Common Technical Specifications 2009/108/EC.

and is in conformity with the national standards transposing harmonised standards:

- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 15223-1:2016
- BS EN ISO 18113-2:2011
- BS EN ISO 23640:2015

The conformity assessment procedure performed was in accordance with Annex IV of Directive 98/79/EC and was carried out by UL International (UK) Ltd, Womersley House, The Guildway, Old Portsmouth Road, Guildford, Surrey GU3 1LR, United Kingdom, Notified Body Number 0843.

The certificates issued by UL-UK Ltd to show compliance are numbers 354.170425 and 355.130523.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 23 May 2017.



Eddy Velthuis
Technical Director



File No A12241;
ISO 13485:2003; ISO 9001:2008

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Email: info@lornelabs.com
www.lornelabs.com

Registered office as above, Registered in England No. 04540797, VAT No. 800 3655 66

EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product Name	Catalogue Number
Anti-B Monoclonal	610010

has been classified as List A (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC) and the Commission Decision on Common Technical Specifications 2009/108/EC.

and is in conformity with the national standards transposing harmonised standards:

- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 15223-1:2016
- BS EN ISO 18113-2:2011
- BS EN ISO 23640:2015

The conformity assessment procedure performed was in accordance with Annex IV of Directive 98/79/EC and was carried out by UL International (UK) Ltd, Womersley House, The Guildway, Old Portsmouth Road, Guildford, Surrey GU3 1LR, United Kingdom, Notified Body Number 0843.

The certificates issued by UL-UK Ltd to show compliance are numbers 354.170425 and 355.130523.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 23 May 2017.



Eddy Velthuis
 Technical Director



File No A12941
 ISO 13485:2003; ISO 9001:2008

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 Berkshire RG6 4UT United Kingdom | www.lornelabs.com

Registered office as above. Registered in England No. 04540797. VAT No. 800 3655 66

EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product Name	Catalogue Number
Anti-D Clone 1 Monoclonal	730010

has been classified as List A (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC) and the Commission Decision on Common Technical Specifications 2009/108/EC.

and is in conformity with the national standards transposing harmonised standards:

- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 15223-1:2016
- BS EN ISO 18113-2:2011
- BS EN ISO 23640:2015

The conformity assessment procedure performed was in accordance with Annex IV of Directive 98/79/EC and was carried out by UL International (UK) Ltd, Womersley House, The Guildway, Old Portsmouth Road, Guildford, Surrey GU3 1LR, United Kingdom, Notified Body Number 0843.

The certificates issued by UL-UK Ltd to show compliance are numbers 354.170425 and 355.130523.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 23 May 2017.



Eddy Velthuis
Technical Director



File No A12241;
ISO 13485:2003; ISO 9001:2008

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Registered office as above. Registered in England No. 04540797. VAT No. 800 3655 66

EC DECLARATION OF CONFORMITY

Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product Name	Catalogue Number
Anti-D Duoclonal Monoclonal	740010

has been classified as List A (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC) and the Commission Decision on Common Technical Specifications 2009/108/EC.

and is in conformity with the national standards transposing harmonised standards:

- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 15223-1:2016
- BS EN ISO 18113-2:2011
- BS EN ISO 23640:2015

The conformity assessment procedure performed was in accordance with Annex IV of Directive 98/79/EC and was carried out by UL International (UK) Ltd, Womersley House, The Guildway, Old Portsmouth Road, Guildford, Surrey GU3 1LR, United Kingdom, Notified Body Number 0843.

The certificates issued by UL-UK Ltd to show compliance are numbers 354.170425 and 355.130523.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 23 May 2017.



Eddy Velthuis
Technical Director



File No A12241:
ISO 13485:2003; ISO 9001:2008

Lorne Laboratories Limited
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www.lornelabs.com

Registered office as above. Registered in England No. 04540797. VAT No. 800 3655 66



Certificat

Certificate

N° 2007/28642.5

AFNOR Certification certifies that the management system implemented by:
AFNOR Certification удостоверяет, что система менеджмента организации:



ZAO "EKOlab"
ЗАО «ЭКОлаб»



for the following activities:
для следующих областей деятельности:

DEVELOPMENT, PRODUCTION, STORAGE AND SALE OF MEDICAL DEVICES FOR IN-VITRO DIAGNOSTICS.

**РАЗРАБОТКА, ПРОИЗВОДСТВО, ХРАНЕНИЕ И РЕАЛИЗАЦИЯ МЕДИЦИНСКИХ ИЗДЕЛИЙ
ДЛЯ IN-VITRO ДИАГНОСТИКИ.**

has been assessed and found to meet the requirements of:
проверена и признана соответствующей требованиям стандарта:

ISO 13485:2016

and is developed on the following locations:
и действует на следующих площадках:

142530, RUSSIA, MOSCOW REGION, ELEKTROGORSK CITY, Budennogo str., 1-1A
142530, РОССИЯ, МОСКОВСКАЯ ОБЛАСТЬ, г. ЭЛЕКТРОГОРСК, ул. Буденного, 1-1А

This certificate is valid from (year/month/day)
Данный сертификат действителен с (год/месяц/день)

2019-06-28

until
до

2022-06-27



Ce document est signé électroniquement. Il constitue un original électronique à valeur probatoire.
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GRUPE 1 - METABOLITES DIVERS
GROUP 1 - MISCELLANEOUS METABOLITES
GRUPO 1 - METABOLICOS VARIOS

DESIGNATION DU REACTIF/ REAGENT DESIGNATION/ DESIGNACION DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO CE	Code GMDN/ GMDN Code/ Codigo GMDN
ALBUMIN	ALBU-0600/0700/0250	DOS-CE-ALBU_R&Sd	53597
ALBUMIN ENVOY	ALBU-0850		53233
BILIRUBIN DIRECT 4+1	BIDI-0600/0250		53279
BILIRUBIN TOTAL 4+1	BITO-0600/0250	DOS-CE-BILI 4+1	53229/53233
BILIRUBIN TOTAL & DIRECT 4+1	BITD-0600		
CREATININE ENVOY	CRSL-0850	DOS-CE-CRSL	53250
CREATININE JAFPE	CRCO-0600/0700	DOS-CE-CRCS_R&Sd	53251
CREATININE PAP SL	CRSL-0630/0250	DOS-CE-CRSL	53250
DIRECT BILIRUBIN ENVOY	BIDV-0850	DOS-CE-BIDV	53233
GLUCOSE ENVOY	GPST-0850	DOS-CE-GPST_R&Sd	
GLUCOSE HK SL	GHSI-0600/0250	DOS-CE-GHSI	53301
GLUCOSE PAP SL	GPST-0495/0500/0700/ 0507/0707/0250/0455/0497	DOS-CE-GPST_R&Sd	
HEMOGLOBIN	HEMO-0400	DOS-CE-HEMO	32430
IRON TIBC	FECA-0850	DOS-CE-TIBC	53904
LACTATE	LACT-0100	DOS-CE-LACT	53342
MICROPROTEIN PLUS	PRTU-0600/0250	DOS-CE-PRTU_R&Sd	53481
PHOSPHORUS	PHOS-0600/0230		
PHOSPHORUS ENVOY	PHOS-0850	DOS-CE-PHOS_R&Sd	59123
TOTAL BILIRUBIN ENVOY	BITV-0850	DOS-CE-BITV	53229
TOTAL PROTEIN	PRTB-0600	DOS-CE-PRTB	
TOTAL PROTEIN ENVOY	PROB-0850		
TOTAL PROTEIN PLUS	PROB-0600/0700/0250	DOS-CE-PROB_R&Sd	53985
UREA ENVOY	URSL-0850	DOS-CE-URSL_R&Sd	
UREA UV	URUV-0400	DOS-CE-URUV	53587
UREA UV SL	URSL-0400/0420/0500/ 0407/0427/0507/0250/0455	DOS-CE-URSL_R&Sd	
URIC ACID	ACUR-0200/0400	DOS-CE-ACUR	
URIC ACID ENVOY	AUVD-0850	DOS-CE-AUVD	
URIC ACID MONO SL	AUMJ-0420/0500/0700/ 0427/0497/0507/0707/0250	DOS-CE-AUML_R&Sd	53583
URIC ACID SL	AUSL-0250	DOS-CE-AUSL	

ELITech Clinical Systems SAS

Zone Industrielle
61500 Sées - France
Tél : +33(0)2 33 81 21 00 - Fax : +33(0)2 22 28 77 51
SIRET 318 365 228 00036

Société par actions simplifiée au capital de 1 219 592,14€ - SIREN : 318 365 228 - RCS ALENCON

DCCF-GI - 132 - Mai /May /Mayo 2018

1/2

DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone Industrielle 61500 Sées France, déclarons sous notre seule responsabilité que les réactifs appartenant au groupe 1 «METABOLITES DIVERS», référencés dans la liste ci-jointe, sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic in vitro et au code de la santé publique.

Cette déclaration est basée sur le contenu de chaque dossier technique DOS-CE-XXXX et s'appuie sur la certification de notre système qualité selon la norme NF EN ISO 13485 : 2016 (Certification valable jusqu'au 27 juillet 2020).

(Voir liste ci-jointe).

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 Sées France, hereby certify, under our own responsibility, that the reagents belonging to Group 1 "MISCELLANEOUS METABOLITES", such as listed hereto, conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to in vitro diagnostic medical devices and to the public health code.

This declaration is based on the contents of each DOS-CE-XXXX technical file and is supported by the certification of our quality system according to the standard NF EN ISO 13485 : 2016 (Certification valid until July 27th, 2020).

(See attached list).

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 Sées France, declaramos bajo nuestra única responsabilidad que los reactivos pertenecientes al grupo 1 "METABOLICOS VARIOS", referenciados en la lista adjunta, son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico in vitro y el código de salud pública.

Esta declaración se basa en el contenido de cada DOS-CE-XXXX técnica y está respaldada por la certificación de nuestro sistema de calidad según la norma NF EN ISO 13485 : 2016 (Certificación válida hasta el 27 de Julio 2020).

(Ver lista adjunta)

Sées, le 18 Mai 2018

Valérie GOURDON,
Responsable des Affaires Réglementaires
Regulatory Affairs Manager
Responsable de los Asuntos Reglamentarios

ELITech Clinical Systems SAS

Zone Industrielle
61500 Sées - France
Tél : +33(0)2 33 81 21 00 - Fax : +33(0)2 22 28 77 51
SIRET 318 365 228 00036

Cécile GOUBAULT,
Directeur Général Délégué
Managing Director
Directora General



Société par actions simplifiée au capital de 1 219 592,14€ - SIREN : 318 365 228 - RCS ALENCON

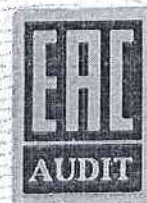
DCCF-GI - 132 - Mai /May /Mayo 2018



ФЕДЕРАЛЬНОЕ АГЕНТСТВО
ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ
СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ ГОСТ Р
«EAC AUDIT»
РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028.04EAC1
ОРГАН ПО СЕРТИФИКАЦИИ ООО «ГОРТЕСТ»
РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028
ИНН 7717616798 ОГРН 1087746489060

Юридический адрес: 109028, Россия, г. Москва, Серебряническая набережная, д. 27,
этаж 4, пом. 1, ком. 17

Телефон: 8 (800) 1000-730, e-mail: info@eacaudit.ru



№003749

СЕРТИФИКАТ СООТВЕТСТВИЯ

Регистрационный номер № 04EAC1.CM.00813

Общество с ограниченной ответственностью «МиниМед»

(наименование лица)

241520, Россия, Брянская область, Брянский район, с. Супонево, ул. Шосейная, д.17А

(юридический адрес лица)

241520, Россия, Брянская область, Брянский район, с. Супонево, ул. Шосейная, д.17А

(фактический адрес лица)

ИНН: 3234007127

ОГРН: 1023202138332

НАСТОЯЩИЙ СЕРТИФИКАТ УДОСТОВЕРЯЕТ СООТВЕТСТВИЕ

системы менеджмента качества изделий медицинских «МиниМед» требованиям ГОСТ ISO 13485-2017 (ISO 13485:2016) «Изделия медицинские. Системы менеджмента качества. Системные требования для целей регулирования» применительно к Производство лабораторной посуды, медицинских изделий, приборов и принадлежностей, красителей, реагентов и наборов реагентов для in-vitro диагностики

Дата регистрации: 19-03-2019

Срок действия до: 18-03-2022

Руководитель органа
по сертификации:



В. И. Погодин

(подпись)

В. И. Погодин

Е. Д. Курбатова

(подпись)

Е. Д. Курбатова



НАСТОЯЩИЙ СЕРТИФИКАТ ОБЯЗЫВАЕТ ОРГАНИЗАЦИЮ ПОДДЕРЖИВАТЬ СОСТОЯНИЕ ВЫПОЛНЯЕМЫХ РАБОТ В СООТВЕТСТВИИ С
ВЫШЕУКАЗАННЫМИ СТАНДАРТАМИ, ЧТО БУДЕТ НАХОДИТЬСЯ ПОД КОНТРОЛЕМ ОРГАНА ПО СЕРТИФИКАЦИИ СИСТЕМЫ
ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ «EAC AUDIT» И ПОДТВЕРЖДАТЬСЯ ПРИ ПРОХОЖДЕНИИ ЕЖЕГОДНОГО ИНСПЕКЦИОННОГО КОНТРОЛЯ

Федеральное агентство по техническому регулированию и метрологии



Система добровольной сертификации "НОПСС". РОСС RU.31827.04ЖСН1

Орган по сертификации ООО "Невский Альянс". ОГРН 1147847286960 ИНН 7842525530

www.nopss.ru

СЕРТИФИКАТ СООТВЕТСТВИЯ

выдан

Общество с ограниченной ответственностью
«МиниМед»

ИНН 3234007127 / ОГРН 1023202138332

241520, Брянская область, Брянский район, с. Спасское, ул. Шоссейная, д. 17А

Подтверждает, что система менеджмента качества
соответствует требованиям ГОСТ ISO 9001-2015 (ISO 9001:2015)

При осуществлении работ согласно приложению №1 к настоящему сертификату

Сертификат выдан на основании решения экспертной комиссии

от 24.09.2018

Срок действия до 24 сентября 2021

Номер в едином реестре системы С1256

Руководитель органа
по сертификации:



Подпись

Платонов Б.А.

Настоящий сертификат обязывает организацию поддерживать состояние выполняемых работ в соответствии с вышеуказанным стандартом, что будет находиться под контролем органа по сертификации СДС "НОПСС" и подтверждаться при прохождении ежегодного инспекционного контроля.

Федеральное агентство по техническому регулированию и метрологии



Система добровольной сертификации "НОПСС". РОСС RU.31827.04ЖСН1

Орган по сертификации ООО "Невский Альянс". ОГРН 1147847286960 ИНН 7842525530

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ПРИЛОЖЕНИЕ №1

К сертификату соответствия № С1256

Применительно к видам деятельности:

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НОПСС



Руководитель органа
по сертификации:

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Регистрационное удостоверение № ФСР 2009/05559 от 04.12.2015 г.

Паспорт

Набор реагентов «Масло иммерсионное» по ТУ 9398-011-29508133-2009

Серия	31-19	Дата изготовления	22.10.2019
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1. Назначение

Используется для ахроматических и апохроматических объективов микроскопов всех видов, кроме люминесцентных, предназначенных для работы в видимой области спектра.

2. Технические требования

Наименование показателя	Характеристика и норма по ТУ	Результаты анализа
Внешний вид	Прозрачная бесцветная жидкость со слабым желтоватым оттенком	соответствует
Вязкость кинематическая при температуре 20°C, мм ² /с	От 220	1317
Показатель преломления при температуре 20°C	От 1,5150 до 1,5180	1,5155
Коэффициент пропускания масла, %	Не менее 70	440 нм – 98,2 540 нм – 100,0

Иммерсионное масло легко удаляется с поверхности препарата, фронтальной линзы и оправы объектива; инертно к окрашенным и неокрашенным препаратам.

Упаковка – флакон-капельница вместимостью 100,0 мл обеспечивает аккуратное и экономичное нанесение масла на препарат.

Срок годности – 1,5 года с даты изготовления.

3. Транспортирование и хранение

Транспортирование должно проводиться всеми видами крытого транспорта в соответствии с правилами перевозки грузов, действующими на данном виде транспорта. Хранение – в упаковке предприятия-изготовителя в прохладном месте при относительной влажности воздуха не более 80% в местах, защищенных от воздействия прямых солнечных лучей, атмосферных осадков и агрессивных сред в течение всего срока годности.

4. Гарантии изготовителя

Изготовитель гарантирует соответствие качества набора реагентов «Масло иммерсионное» требованиям ТУ 9398-011-29508133-2009 при соблюдении потребителем условий транспортирования, хранения и применения в течение всего срока годности.

Начальник ПТО



Бабич В.А.





Italia

CERTIFICATO

Nr. 50 100 11497 - Rev. 003

Si attesta che / This is to certify that

IL SISTEMA QUALITÀ DI
THE QUALITY SYSTEM OF**LIOFILCHEM S.r.l.**SEDE LEGALE E OPERATIVA:
REGISTERED OFFICE AND OPERATIONAL SITE:**VIA SCOZIA SNC - ZONA INDUSTRIALE
I-64026 ROSETO DEGLI ABRUZZI (TE)**SEDE OPERATIVA:
OPERATIONAL SITE:**CONTRADA PIANE VOMANO – TRAVERSA DI VIA GRECIA
I-64026 ROSETO DEGLI ABRUZZI (TE)**È CONFORME AI REQUISITI DELLA NORMA
HAS BEEN FOUND TO COMPLY WITH THE REQUIREMENTS OF**UNI EN ISO 9001:2015**QUESTO CERTIFICATO È VALIDO PER IL SEGUENTE CAMPO DI APPLICAZIONE
THIS CERTIFICATE IS VALID FOR THE FOLLOWING SCOPE**Progettazione e sviluppo, produzione e commercializzazione di
dispositivi medico diagnostici in-vitro: terreni di coltura per
batteriologia, sistemi di identificazione e antibiogramma, kit per la
determinazione di plasmaproteine (IAF 12, 29)****Design and development, production and sale of in-vitro diagnostic
medical devices: culture media for bacteriology, identification and
susceptibility testing systems, kits for plasma protein determination
(IAF 12, 29)**

SGQ N° 049A

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For the Certification Body
TÜV Italia S.r.l.

Validità / Validity

Dal / From: 2019-02-11

Al / To: 2022-02-10

Data emissione / Issuing Date

Andrea Coscia

Direttore Divisione Business Assurance

2019-02-11

PRIMA CERTIFICAZIONE / FIRST CERTIFICATION: 2012-09-25LA VALIDITÀ DEL PRESENTE CERTIFICATO È SUBORDINATA A SORVEGLIANZA PERIODICA A 12 MESI E AL RIESAME COMPLETO DEL SISTEMA DI
GESTIONE AZIENDALE CON PERIODICITÀ TRIENNALE"THE VALIDITY OF THE PRESENT CERTIFICATE DEPENDS ON THE ANNUAL SURVEILLANCE EVERY 12 MONTHS AND ON THE COMPLETE REVIEW OF
COMPANY'S MANAGEMENT SYSTEM AFTER THREE-YEARS"

CERTIFICATO

N° Q5 071067 0006 Rev. 00

Titolare del certificato:

Liofilchem S.r.l.

Via Scozia
64026 Roseto degli Abruzzi (TE)
ITALIA

Stabilimento(i):

Liofilchem S.r.l.
Via Scozia, 64026 Roseto degli Abruzzi (TE), ITALIA
Liofilchem S.r.l.
Contrada Piane Vomano, Traversa di Via Grecia,
64026 Roseto degli Abruzzi (TE), ITALIA

**Marchio di
certificazione:**



Campo di applicazione:

Progettazione e sviluppo, produzione e commercializzazione di dispositivi medico diagnostici In-vitro: terreni di coltura per batteriologia, sistemi di identificazione e antibiogramma, kit per la determinazione di plasmaproteine. Distribuzione di altri dispositivi medico diagnostici in-vitro

Norma(e) applicata(e):

EN ISO 13485:2016
Dispositivi medici – Sistemi di gestione per la qualità -
Requisiti per scopi regolamentari
(ISO 13485:2016)
DIN EN ISO 13485:2016

L'Organismo di Certificazione TÜV SÜD Product Service GmbH certifica che la società sopramenzionata ha istituito e mantiene un sistema di gestione qualità conforme ai requisiti della(e) norma(e) elencata(e). Vedere anche note sul retro.

N° del rapporto:

ITA1070742

**Valido da:
Valido fino al:**

2018-12-19
2021-12-18

Data, 2018-12-19

S. Preiß

Stefan Preiß



Pagina 1 di 1

Traduzione per scopi informativi. La sola versione inglese (tedesca) è legalmente impegnativa.

GRAM COLOR KIT

DESCRIZIONE

GRAM COLOR KIT è un kit per la colorazione dei microrganismi, che ne permette la differenziazione in due categorie: Gram-positivi (Gram+) che si colorano in blu e Gram-negativi (Gram-), che si colorano in rosso. Tale colorazione costituisce, insieme con l'osservazione diretta della morfologia cellulare, il primo livello di classificazione tassonomica dei procarioti.

CONTENUTO DELLE CONFEZIONI

I reagenti sono contenuti in flaconi di plastica, chiusi in termoisolamento e forniti di tappo goccia. Ciascuna confezione contiene:

- 1 flacone contenente 250 ml di Soluzione Cristal Violetto
- 1 flacone contenente 250 ml di Soluzione Lugol-PVP
- 1 flacone contenente 250 ml di Soluzione Decolorante
- 1 flacone contenente 250 ml di Soluzione Safranina
- 1 foglio istruzioni

PRINCIPIO DEL METODO

La colorazione di Gram è basata sulla proprietà che ha il Cristal Violetto di combinarsi con lo iodio, formando composti non decolorabili con l'alcool o con la miscela alcool-acetone. Alcuni batteri hanno una speciale affinità per questa reazione e, una volta colorati con il cristal violetto, non perdono il colore, se trattati con l'alcool o con la miscela alcool-acetone, restando colorati in blu (batteri Gram-positivi). Altri perdono il colore blu e si colorano con la Safranina assumendo una colorazione rossa (batteri Gram-negativi).

COMPOSIZIONE

Le soluzioni che compongono il GRAM COLOR KIT sono tutte soluzioni acquose ad eccezione della Soluzione Decolorante.

Tabella n°1

GRAM COLOR KIT			
Contenuto			
Soluzione Cristal Violetto	1	Cristal Violetto	2%
		Alcool etilico	20%
		Ossalato d'ammonio	0.8%
		Iodio	1.3%
Soluzione Lugol-PVP	2	Ioduro di potassio	2%
		PVP (Polivinilpirrolidone)	7%
		Alcool 95°C	50%
Soluzione Decolorante	3	Acetone	50%
		Safranina	0.25%
Soluzione Safranina	4	Alcool etilico	10%

RACCOLTA DEI CAMPIONI

I campioni da sottoporre alla colorazione di Gram sono costituiti principalmente da materiale clinico e da colture microbiche.

Le colonie da sottoporre alla colorazione di Gram devono essere prelevate da colture giovani (18-24 ore) preferibilmente da terreni agarizzati.

PROCEDURA DEL TEST

Preparazione e fissazione

Utilizzando vetrini puliti, eseguire uno striscio della coltura o del materiale patologico. Lasciare essiccare all'aria e fissare al calore con passaggi rapidi sulla fiamma. Eseguire la fissazione del campione evitando un eccesso di riscaldamento. Si possono adottare anche altri metodi di fissazione.

Colorazione

- Ricoprire il vetrino con la Soluzione Cristal Violetto. Attendere 1 minuto, quindi lavare delicatamente con acqua.
- Ricoprire il vetrino con la Soluzione Lugol-PVP. Attendere 1 minuto, quindi lavare delicatamente con acqua.
- Decolorare con la Soluzione Decolorante finché il preparato libera colorante (circa 30-60 secondi), quindi lavare delicatamente con acqua.
- Ricoprire il vetrino con la Soluzione Safranina. Attendere 30-60 secondi, quindi lavare delicatamente con acqua.
- Asciugare.
- Osservare il preparato al microscopio con obiettivo per immersione.

INTERPRETAZIONE DEI RISULTATI

I microrganismi Gram-negativi appaiono di colore rosso.

I microrganismi Gram-positivi appaiono di colore blu.

La colorazione di Gram permette di differenziare:

- I bacilli Gram-negativi da quelli Gram-positivi;
- I cocci Gram-negativi da quelli Gram-positivi;
- I coccobacilli Gram-negativi da quelli Gram-positivi; i diplococchi Gram-negativi da quelli Gram-positivi.

CONTROLLO QUALITÀ

Ogni lotto di GRAM COLOR KIT viene sottoposto al controllo di qualità utilizzando una coltura di *Escherichia coli* ATCC 25922 per il controllo dei batteri Gram-negativi (colore rosso) ed una coltura di *Staphylococcus aureus* ATCC 25923 per il controllo dei batteri Gram-positivi (colore blu).

LIMITI

1. La colorazione di Gram fornisce una preliminare identificazione ma non sostituisce i normali studi culturali del campione.
2. Terapie antibiotiche possono rendere i batteri gram-positivi più sensibili alla decolorazione ed apparire di colore rosa-rosso invece di blu.
3. Le cellule prelevate da colture giovani di 18-24 ore hanno una maggiore affinità per i coloranti rispetto alle cellule prelevate da colture vecchie.
4. La colorazione di Gram viene alterata dalla distruzione fisica della parete cellulare o del protoplasma; infatti la parete cellulare dei batteri Gram-positivi interpone una barriera che impedisce il rilascio del complesso Cristal Violetto-iodio dal citoplasma e la parete cellulare dei batteri Gram-negativi contiene lipidi solubili in solventi organici che permettono la decolorazione del citoplasma. Pertanto i microrganismi distrutti fisicamente da un eccesso di calore non reagiscono alla colorazione di Gram come atteso.

PRECAUZIONI

La confezione di GRAM COLOR KIT contiene sostanze classificate come pericolose ai sensi della legislazione vigente; per il suo impiego si consiglia di consultare la scheda di sicurezza.

GRAM COLOR KIT è un kit per la colorazione batterica, da usare solo per uso diagnostico *in vitro*, è destinato ad un ambito professionale e deve essere usato in laboratorio da operatori adeguatamente addestrati, con metodi approvati di asepsi e di sicurezza nei confronti degli agenti patogeni.

CONSERVAZIONE

Conservare GRAM COLOR KIT a 10-25°C nella sua confezione originale. Non conservare vicino a fonti di calore ed evitare eccessive variazioni di temperatura. In queste condizioni il prodotto GRAM COLOR KIT è valido fino alla data di scadenza indicata in etichetta. Non utilizzare oltre questa data. Eliminare se vi sono segni di deterioramento (cambiamenti di colore delle soluzioni o presenza di precipitati grossolani).

ELIMINAZIONE DEL MATERIALE USATO

Dopo l'utilizzazione, i vetrini colorati con il GRAM COLOR KIT ed il materiale venuto a contatto con il campione devono essere decontaminati e smaltiti in accordo con le tecniche in uso in laboratorio per la decontaminazione e lo smaltimento di materiale potenzialmente infetto.

BIBLIOGRAFIA

- Kruczak-Filipov, P., and R.G. Shively. 1992. Gram stain procedure, p.1.5.1-1.5.18. In H.D. Isenberg (ed.) Clinical Microbiology Procedures Handbook, vol. 1. American Society for Microbiology, Washington, D.C.
- Murray, P.R. (ed.) 1999. Manual of Clinical Microbiology, 7th ed. American Society of Microbiology, Washington, D.C.

PRESENTAZIONE

Prodotto		
GRAM COLOR KIT	80293	4 x 250 ml

TABELLA DEI SIMBOLI

IVD	Dispositivo medico diagnostico <i>in vitro</i>		Non riutilizzare
	Fabbricante		Contenuto sufficiente per <n> saggi
REF	Numero di catalogo		Fragile, maneggiare con cura
	Utilizzare entro		Attenzione, vedere le istruzioni per l'uso
	Limiti di temperatura	LOT	Codice del lotto



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F00411

Rev.2/ 13.09.2011



ZIEHL NEELSEN KIT

ITALIANO

Kit per la colorazione dei batteri alcool-acido resistenti

DESCRIZIONE

ZIEHL NEELSEN KIT è un kit per la colorazione dei batteri alcool-acido resistenti presenti in campioni patologici o in terreni culturali. Questa colorazione è particolarmente utile per eseguire l'esame microscopico per la ricerca dei micobatteri.

CONTENUTO DELLE CONFEZIONI

I reagenti sono contenuti in flaconi di plastica, chiusi in termoisolamento e forniti di tappo goccia.

Ciascuna confezione contiene:

- 1 flacone contenente 250 ml di Soluzione Fucsina
- 1 flacone contenente 250 ml di Soluzione Alcool-Acido
- 1 flacone contenente 250 ml di Soluzione Blu di Metilene
- 1 foglio istruzioni

PRINCIPIO DEL METODO

La colorazione di ZIEHL NEELSEN è basata sulla capacità del fenolo di permettere la penetrazione della Fucsina basica all'interno della cellula batterica. Alcuni batteri, una volta colorati con la Fucsina basica, se trattati con la Soluzione Alcool-Acido, restano colorati in rosso (batteri alcool-acido resistenti). Altri perdono il colore rosso e si colorano con il Blu di Metilene assumendo una colorazione blu (batteri sensibili alla decolorazione con alcool-acido).

COMPOSIZIONE

La formulazione dei costituenti dello ZIEHL NEELSEN KIT è quella prevista per la colorazione dei micobatteri. Le tre soluzioni che ne fanno parte sono tutte soluzioni acquose ad eccezione della Soluzione Alcool-Acido.

Tabella n°1

	Ziehl Neelsen Kit		
	Contenuto		
1	Soluzione Fucsina	Fucsina basica Alcool etilico 95% Fenolo	0.3% 10% 5%
2	Soluzione Alcool-Acido	Alcool etilico 95% HCl concentrato	97% 3%
3	Soluzione Blu di Metilene	Blu di Metilene	0.3%

RACCOLTA DEI CAMPIONI

I campioni da sottoporre alla colorazione degli alcool-acido resistenti sono costituiti principalmente da materiale clinico e da colture microbiche.

PROCEDURA DEL TEST

Preparazione e fissazione

Utilizzando vetrini puliti, eseguire lo striscio del materiale patologico o della coltura. Lasciare essiccare all'aria e fissare al calore con passaggio rapido sulla fiamma o con altri metodi di fissazione. Eseguire la fissazione del campione evitando un eccesso di riscaldamento.

Colorazione

I lavaggi del preparato, in tutte le fasi della colorazione, devono essere fatti con acqua priva di specie alcool-acido-resistenti.

- Ricoprire il vetrino con la Soluzione Fucsina. Scaldare lentamente il vetrino fino alla formazione di vapori. Mantenere il riscaldamento del vetrino per 3-5 minuti, lasciare raffreddare e lavare quindi con acqua.
- Decolorare con la Soluzione Alcool-Acido finché il preparato libera colorante (circa 1-2 minuti). Lavare con acqua.
- Ricoprire il vetrino con la Soluzione Blu di Metilene. Lasciare a contatto per 30-60 secondi quindi lavare con acqua e lasciare asciugare.
- Osservare il preparato al microscopio con obiettivo per immersione.

INTERPRETAZIONE DEI RISULTATI

I micobatteri e i batteri alcool-acido-resistenti appaiono colorati in rosso mentre il materiale di fondo ed altri batteri non alcool-acido resistenti appaiono colorati in blu.

CONTROLLO QUALITÀ

Ogni lotto di ZIEHL NEELSEN KIT viene sottoposto al controllo di qualità utilizzando una coltura di *Mycobacterium smegmatis* ATCC 14468 per il controllo dei batteri alcool-acido resistenti (colore rosso) ed una coltura di *Staphylococcus aureus* ATCC 25923 per il controllo dei batteri sensibili alla decolorazione con alcool-acido (colore blu).

LIMITI

- La colorazione di ZIEHL NEELSEN fornisce una presuntiva evidenza della presenza di microrganismi alcool-acido resistenti nel campione, evidenza che deve essere confermata con altri metodi. Un risultato negativo della colorazione non indica necessariamente che il campione sia negativo per la presenza di microrganismi alcool-acido resistenti.
- I micobatteri a rapida crescita possono trattenere i coloranti a vari livelli.
- Quando si decolora con la Soluzione Alcool-Acido, evitare una decolorazione insufficiente.

PRECAUZIONI

La confezione di ZIEHL NEELSEN KIT contiene sostanze classificate come pericolose ai sensi della legislazione vigente; per il suo impiego si consiglia di consultare la scheda di sicurezza. ZIEHL NEELSEN KIT è un kit per la colorazione batterica da usare solo per uso diagnostico *in vitro*, è destinato ad un ambito professionale e deve essere usato in laboratorio da operatori adeguatamente addestrati, con metodi approvati di asepsi e di sicurezza nei confronti degli agenti patogeni.

CONSERVAZIONE

Conservare ZIEHL NEELSEN KIT a 10-25°C nella sua confezione originale. Non conservare vicino a fonti di calore ed evitare eccessive variazioni di temperatura. In queste condizioni il prodotto ZIEHL NEELSEN KIT è valido fino alla data di scadenza indicata in etichetta. Non utilizzare oltre questa data. Eliminare se vi sono segni di deterioramento (cambiamenti di colore delle soluzioni o presenza di precipitati grossolani).

ELIMINAZIONE DEL MATERIALE USATO

Dopo l'utilizzazione, i vetrini colorati con il ZIEHL NEELSEN KIT ed il materiale venuto a contatto con il campione devono essere decontaminati e smaltiti in accordo con le tecniche in uso in laboratorio per la decontaminazione e lo smaltimento di materiale potenzialmente infetto.

PRESENTAZIONE

Prodotto	REF	
ZIEHL NEELSEN KIT	80276	3 x 250 ml

TABELLA DEI SIMBOLI

IVD Dispositivo medico diagnostico <i>in vitro</i>	Non riutilizzare	Fabbricante	Contenuto sufficiente per <n> saggi	Limiti di temperatura
REF Numero di catalogo	Fragile, maneggiare con cura	Utilizzare entro	Attenzione, vedere le istruzioni per l'uso	LOT Codice del lotto