



Certificate

No. Q5 020747 0242 Rev. 02

Holder of Certificate: **Nova Biomedical Corporation**
200 Prospect Street
Waltham MA 02454
USA

Certification Mark:



Scope of Certificate: Design and Development, Production, Distribution, Installation, Servicing and Technical Support of In-Vitro Diagnostic Reagents (Calibrators, Controls, Reagents, Sensors and Test Cartridges) and Instruments for Clinical Chemistry, Blood Gas and Hematology, including Near Patient / Point of Care and Self-Testing devices; The provision of manufacturing services of In-Vitro Diagnostic Reagents (Calibrators, Controls) for Clinical Chemistry, Blood Gas and Hematology, In-Vitro Diagnostic General Use Consumables; and Distribution of Lancets.

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with. For details and certificate validity see: [www.tuvsud.com/ps-cert?q=cert:Q5 020747 0242 Rev. 02](http://www.tuvsud.com/ps-cert?q=cert:Q5_020747_0242_Rev_02)

Report No.: 72198686

Valid from: 2024-10-25
Valid until: 2027-10-24

Date, 2024-10-04

Christoph Dicks
Head of Certification/Notified Body

Certificate

No. Q5 020747 0242 Rev. 02

Applied Standard(s): ISO 13485:2016
(EN ISO 13485:2016/AC:2018, EN ISO 13485:2016/A11:2021)
Medical devices - Quality management systems -
Requirements for regulatory purposes

Facility(ies): **Nova Biomedical Corporation**
200 Prospect Street, Waltham MA 02454, USA

Design and Development, Production, Distribution, Installation,
Servicing and Technical Support of In-Vitro Diagnostic Reagents
(Calibrators, Controls, Reagents, Sensors and Test Cartridges)
and Instruments for Clinical Chemistry, Blood Gas and
Hematology, including Near Patient / Point of Care and Self-
Testing devices; the provision of manufacturing services of In-Vitro
Diagnostic Reagents (Calibrators, Controls) for Clinical Chemistry,
Blood Gas and Hematology and In-Vitro Diagnostic General Use
Consumables.

Nova Biomedical Corporation
39 Manning Road, Billerica MA 01821, USA

Production of Self-Testing and Near Patient / Point of Care test
strips.

Nova Biomedical Corporation
165 Lexington Road, Billerica MA 01821, USA

Production of Self-Testing and Near Patient / Point of Care
Instruments

Nova Biomedical Corporation
4 Enterprise Road, Billerica MA 01821, USA

Production of In-Vitro Diagnostic Instruments including Near
Patient / Point of Care; Distribution of Finished Goods; Distribution
of Lancets.



**Self-Declaration of Conformity To European Parliament and Council Directive 98/79/EC
In Vitro Diagnostic Medical Device Directive (IVDD)**

Product name: Nova Stat Profile Prime Plus Analyzer System including Reagents, Calibrators and Controls

Catalog Numbers: List Attached (Two Pages)

Classification: Other/General

Manufacturer: Nova Biomedical Corporation
200 Prospect Street
Waltham, MA 02454 USA

Representative: William Jacques, Director of Regulatory and Quality

Authorized Representative: Nova Biomedical GmbH
Hessenring 13 A, Geb. G
64546 Mörfelden-Walldorf
Germany
Tel: +49 6105 4505-0

Conformity Assessment Route: Annex III

Nova Biomedical, Inc. declares that the products listed are in conformity with the provisions of the Council Directive 98/79/EC for in vitro diagnostic medical devices, including applicable essential requirements of Annex I for legal application of the CE Mark. All supporting documentation is retained under the premises of the manufacturer.

Nova Biomedical, Inc. declares that the electronic products listed are in conformity with the provisions of the Council Directive 2011/65/EC on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS).

Standards Applied:

EN ISO 13485:2016	Medical devices - Quality management systems - Requirements for regulatory purposes
EN 50581:2012	Technical Documentation for the Assessment of Electrical and Electronic Products with Respect to the Restriction of Hazardous Substances
EN 61010-1:2010	Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements
EN 61010-2:101:2015	Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

Signature:

William Jacques, Director of Regulatory and Quality



Date:

Jul/24/2020

List of Catalog Items Covered:

Catalog Number	Product Name	Global Medical Device Nomenclature (GMDN) Name	GMDN Number	DIMDI EDMS Code
57400	Stat Profile Prime Plus® Analyzer	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
59508	Stat Profile Prime Plus® Analyzer (Remanufactured)	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
57820	Stat Profile Prime Plus MicroSensor Card™ with COOX	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
57821	Stat Profile Prime Plus MicroSensor Card™ BUN, Creatinine	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
57822	Stat Profile Prime Plus MicroSensor Card™ with COOX (High Volume)	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
57823	Stat Profile Prime Plus Reference Cartridge	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
57825	Stat Profile Prime Plus Calibrator Cartridge 100 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57826	Stat Profile Prime Plus Calibrator Cartridge 200 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57827	Stat Profile Prime Plus Calibrator Cartridge 300 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57828	Stat Profile Prime Plus Calibrator Cartridge 400 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57829	Stat Profile Prime Plus Calibrator Cartridge 500 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57831	Stat Profile Prime Plus Calibrator Cartridge 100 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57832	Stat Profile Prime Plus Calibrator Cartridge 200 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57833	Stat Profile Prime Plus Calibrator Cartridge 300 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57834	Stat Profile Prime Plus Calibrator Cartridge 400 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57835	Stat Profile Prime Plus Calibrator Cartridge 500 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, calibrator	52859	11-04-04-03-00
57838	Stat Profile Prime Plus Auto QC Cartridge 160 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57839	Stat Profile Prime Plus Auto QC Cartridge 320 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57840	Stat Profile Prime Plus Auto QC Cartridge 480 Sample	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57841	Stat Profile Prime Plus Auto QC Cartridge 105 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57842	Stat Profile Prime Plus Auto QC Cartridge 210 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57843	Stat Profile Prime Plus Auto QC Cartridge 315 Sample with Creat / BUN	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57844	Stat Profile Prime Plus Ampuled Controls BG, COOX Levels 1, 2, 3	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
57845	Stat Profile Prime Plus Ampuled Controls Chemistry Levels 4,5	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
58379	Stat Profile Prime Plus BUN, Creatinine - Blank Sensor Card	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
58642	Stat Profile Prime Plus MicroSensor Card™	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02
58643	Stat Profile Prime Plus MicroSensor Card™ (High Volume)	Point-of-care single channel clinical chemistry analyzer IVD	56681	21-02

Catalog Number	Product Name	Global Medical Device Nomenclature (GMDN) Name	GMDN Number	DIMDI EDMS Code
55229	Nova Linearity Level 1,2,3,4	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-01-00
56198	Linearity Standard Set G Multipack	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-90-00
61656	Nova Linearity Creatinine/BUN/Hct Levels 1,2,3,4	Multiple blood gas/haemoximetry/electrolyte analyte IVD, control	52860	11-50-90-90-00



Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	Ea IgG CODE: EAG.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 – JUNE 2010
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – MARCH 2019
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE
EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado nº/Certificate no	Fecha de validez/Date of validity	ON nº/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	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: 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 el Anexo de este Certificado/Specified in Annex to this Certificate

Elaborado en/In the facilities:

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

Fecha inicial/ Initial date: 11/12/2003

Fecha de prórroga anterior/ Previous extension date: 26/11/2018

Este certificado debe ir acompañado por certificado de examen de diseño: SI / This certificate must be accompanied by design examination certificate: YES

Este certificado es consecuencia de la auditoria del sistema completo de garantía de calidad y del examen de la documentación técnica contenida en el expediente nº 2003 05 0240, y garantiza que los productos descritos cumplen los requisitos de la Directiva./ This certificate is issued on the full quality assurance system audit, and the examination of the technical documentation contained in dossier nº 2003 05 0240, and guarantees that the described products fulfils the requirements of the Directive.

Madrid, 19 de mayo de 2022

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



agencia española de
medicamentos y
productos sanitarios

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 1 de 7

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	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: Idem

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

Clasificación/ Classification: Lista A del Anexo II / List A of Annex II

1. Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante técnicas de Inmunoabsorción enzimática (ELISA)/ Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Enzyme-linked immunosorbent assay (ELISA) [NANDO: IVD 0201; IVD 0202; IVD 0203]

1.1. HBs Ab

- SAB.CE (96 tests)

Descrito en el certificado/ Described in the certificate 2003 12 0390 ED

1.2. HBc Ab

- BCAB.CE (96 tests)

Descrito en el certificado/ Described in the certificate 2003 12 0391 ED

1.3. HBc IgM

- BCM.CE (96 tests)

Descrito en el certificado/ Described in the certificate 2004 03 0424 ED

1.4. HBe Ag & Ab

- HBE.CE (96 tests)

Descrito en el certificado/ Described in the certificate 2004 03 0425 ED

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 2 de 7

C/ CAMPEZO, 1 - EDIFICIO 8

28022 MADRID

Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97

Fax: (+34) 91.822.52.89

ORGANISMO NOTIFICADO 0318



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	0318

1.5. HBs Ag Confirmation

- SCONF.CE (20 tests) Descrito en el certificado/ Described in the certificate 2006 11 0511 ED
- SCONF.CE.40 (40 tests)

1.6. HBs Ag one Version ULTRA

- SAG1ULTRA.CE (192 tests) Descrito en el certificado/ Described in the certificate 2008 12 0588 ED
- SAG1ULTRA.CE.96 (96 tests)
- SAG1ULTRA.CE.480 (480 tests)
- SAG1ULTRA.CE.960 (960 tests)
- SAG1ULTRA.CE.DB (192 tests)

1.7. HCV Ab

- CVAB.CE (192 tests) Descrito en el certificado/ Described in the certificate 2003 12 0392 ED
- CVAB.CE.96 (96 tests)
- CVAB.CE.480 (480 tests)
- CVAB.CE.960 (960 tests)
- CVAB.CE.DB (192 tests)

1.8. HCV Ab Confirmation

- CCONF.CE (12 tests) Descrito en el certificado/ Described in the certificate 2005 09 0485 ED

1.9. HCV IgM

- CVM.CE (96 tests) Descrito en el certificado/ Described in the certificate 2007 09 0532 ED

MODELO -1 ANEXO IV CT Cert. 98/79/1-Rev. -18/05/2020

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 3 de 7

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID

Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89

ORGANISMO NOTIFICADO 0318



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	0318

1.10. HCV Ab (Format 20)

- CVAB.CE.EG (192 tests)
- CVAB.CE.EG.96 (96 tests)
- CVAB.CE.EG.480 (480 tests)
- CVAB.CE.EG.960 (960 tests)

Descrito en el certificado/ Described in the
certificate 2015 10 0842 ED

1.11. HDV Ab

- DAB.CE (96 tests)

Descrito en el certificado/ Described in the
certificate 2003 12 0393 ED

1.12. HDV Ag

- DAG.CE (96 tests)

Descrito en el certificado/ Described in the
certificate 2003 12 0394 ED

1.13. HDV IgM

- DIM.CE (96 tests)

Descrito en el certificado/ Described in the
certificate 2003 12 0395 ED

1.14. HTLV I & II Ab Version ULTRA

- HTLVABULTRA.CE (192 tests)
- HTLVABULTRA.CE.96 (96 tests)
- HTLVABULTRA.CE.480 (480 tests)
- HTLVABULTRA.CE.960 (960 tests)
- HTLVABULTRA.CE.DB (192 tests)

Descrito en el certificado/ Described in the
certificate 2011 11 0775 ED

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 4 de 7

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
2003 12 0388 CT	Desde/From	Hasta/To	0318
	20-05-2022	26-05-2025	

1.15. HIV Ab & Ag

- IVCOMB.CE (192 tests)
- IVCOMB.CE.96 (96 tests)
- IVCOMB.CE.480 (480 tests)
- IVCOMB.CE.960 (960 tests)
- IVCOMB.CE.DB (192 tests)

Descrito en el certificado/ *Described in the certificate* 2008 02 0539 ED

2. Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante técnicas de PCR en tiempo real/ Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Real-Time PCR [NANDO: IVD 0203]

2.1 HBV DNA Quantitation (QT)

- HBVDNAQT.CE (50 tests)
- HBVDNAQT.CE.25 (25 tests)
- HBVDNAQT.CE.100 (100 tests)
- HBVDNAQT.CE.150 (150 tests)

Descrito en el certificado/ *Described in the certificate* 2012 09 0790 ED

2.2 HDV RNA Quantitation (QT)

- DRNA.CE (50 tests)
- DRNA.CE.25 (25 tests)
- DRNA.CE.100 (100 tests)
- DRNA.CE.150 (150 tests)

Descrito en el certificado/ *Described in the certificate* 2009 11 0660 ED

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 5 de 7

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	0318

2.3 HDV ONESTEP Quantitation (QT)

- HDVONEQT.CE (50 tests) Descrito en el certificado/ Described in the certificate 2022 04 0973 ED
- HDVONEQT.CE.25 (25 tests)
- HDVONEQT.CE.100 (100 tests)

3 Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante ensayos de quimioluminiscencia (CLIA)/ Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Chemiluminescence Immunoassay (CLIA) [NANDO: IVD 0201; IVD 0202; IVD 0203]

3.1 DIA.CHEMILUX HCV Ab

- RACVAB.CE (100 tests) Descrito en el certificado/ Described in the certificate 2015 01 0834 ED

3.2 DIA.CHEMILUX HBs Ag

- RASAG.CE (100 tests) Descrito en el certificado/ Described in the certificate 2015 10 0841 ED

3.3 DIA.CHEMILUX HIV Ab & Ag

- RAIVCOMB.CE (100 tests) Descrito en el certificado/ Described in the certificate 2016 02 0844 ED

3.4 DIA.CHEMILUX HBc Ab

- RABCAB.CE (100 tests) Descrito en el certificado/ Described in the certificate 2017 07 0863 ED

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 6 de 7

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

**CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE**

**EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC**

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2003 12 0388 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	0318

3.5 DIA.CHEMILUX HTLV I & II Ab

- RAHTLVAB.CE (100 tests) Descrito en el certificado/ Described in the certificate 2018 11 0878 ED

3.6 DIA.CHEMILUX HDV Ab

- RADAB.CE (100 tests) Descrito en el certificado/ Described in the certificate 2020 07 0932 ED

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 mayo de 2022

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

 **agencia española de
medicamentos y
productos sanitarios**

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: NDASRWLD3A



CORREO ELECTRÓNICO
on0318@aemps.es

Página 7 de 7

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE
EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2004 05 0442 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	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: 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 el Anexo de este Certificado/Specified in Annex to this Certificate

Elaborado en/In the facilities:

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

Fecha inicial/ Initial date: 10/05/2014

Fecha de prórroga anterior/ Previous extension date: 26/11/2018

Este certificado debe ir acompañado por certificado de examen de diseño: NO / This certificate must be accompanied by design examination certificate: NO

Este certificado es consecuencia de la auditoria del sistema completo de garantía de calidad y del examen de la documentación técnica contenida en el expediente n° 2003 05 0240, y garantiza que los productos descritos cumplen los requisitos de la Directiva./ This certificate is issued on the full quality assurance system audit, and the examination of the technical documentation contained in dossier n° 2003 05 0240, and guarantees that the described products fulfils the requirements of the Directive.

Madrid, 19 de mayo de 2022

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



agencia española de
medicamentos y
productos sanitarios

Fdo. Mª Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 1 de 6

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity	ON n°/NB no
2004 05 0442 CT	Desde/From 20-05-2022 Hasta/To 26-05-2025	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: Idem

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

Clasificación/ Classification: Lista B del Anexo II / *List B of Annex II*

1. Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante técnicas de Inmunoabsorción enzimática (ELISA)/ *Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Enzyme-linked immunosorbent assay (ELISA)* [NANDO: IVD 0303; IVD 0305]

1.1. CMV IgM

- CMV.CE (96 tests)

1.2. CMV IgG

- CMVG.CE (96 tests)

1.3. Toxo IgM

- TOXOM.CE (96 tests)

1.4. Toxo IgG

- TOXOG.CE (96 tests)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 2 de 6

C/ CAMPEZO, 1 - EDIFICIO 8

28022 MADRID

Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97

Fax: (+34) 91.822.52.89

ORGANISMO NOTIFICADO 0318



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
2004 05 0442 CT	Desde/From	Hasta/To	0318
	20-05-2022	26-05-2025	

1.5. RUB IgM

- RUBM.CE (96 tests)

1.6. RUB IgG

- RUBG.CE (96 tests)
- RUBG.CE.192 (192 tests)
- RUBG.CE.480 (480 tests)

1.7. TORCH IgM

- TORCHM.CE (96 tests)

1.8. Chlamydia Trachomatis IgG

- CTG.CE (96 tests)

1.9. Chlamydia Trachomatis IgM

- CTM.CE (96 tests)

1.10. Chlamydia Trachomatis IgA

- CTA.CE (96 tests)

1.11. Chlamydia Pneumoniae IgG

- CPG.CE (96 tests)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 3 de 6

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

**EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC**

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
2004 05 0442 CT	Desde/From	Hasta/To	0318
	20-05-2022	26-05-2025	

1.12. Chlamydia Pneumoniae IgM

- CPM.CE (96 tests)

1.13. Chlamydia Pneumoniae IgA

- CPA.CE (96 tests)

2. Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante técnicas de PCR en tiempo real/ Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Real-Time PCR [NANDO: IVD 0303; IVD 0305]

2.1. CMV DNA Quantitation (QT) 2nd Generation

- CMVDNAQT.2G. CE (50 tests)
- CMVDNAQT.2G.CE.25 (25 tests)
- CMVDNAQT.2G.CE.100 (100 tests)
- CMVDNAQT.2G.CE.150 (150 tests)

2.2. Dx CMV Assay

- 37020 (96 tests)

2.3. Toxoplasma Gondii DNA

- TOXODNA.CE (50 tests)
- TOXODNA.CE.25 (25 tests)
- TOXODNA.CE.100 (100 tests)
- TOXODNA.CE.150 (150 tests)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 4 de 6

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

**CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE**

**EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC**

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
2004 05 0442 CT	Desde/From	Hasta/To	0318
	20-05-2022	26-05-2025	

2.4. Chlamydia Trachomatis DNA

- CTDNA.CE (50 tests)
- CTDNA.CE.25 (25 tests)
- CTDNA.CE.100 (100 tests)
- CTDNA.CE.150 (150 tests)

2.5. PRIME MDx CMV DNA Quantitative detection kit

- 56449 (24 tests)
- 56450 (48 tests)

2.6. PRIME MDx Toxoplasma gondii DNA detection kit

- 5647 (24 tests)
- 5648 (48 tests)

3. Reactivos y productos reactivos para la determinación, confirmación y cuantificación de marcadores de infección en muestras humanas mediante ensayos de quimioluminiscencia (CLIA)/ Reagents and reactive products for the determination, confirmation and quantification of infection markers in human samples by Chemiluminescence Immunoassay (CLIA) [NANDO: IVD 0303; IVD 0305]

3.1 DIA.CHEMILUX Cytomegalovirus IgM

- RACMVM.CE (100 tests)

3.2 DIA.CHEMILUX Cytomegalovirus IgG

- RACMVG.CE (100 tests)

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 5 de 6

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89



ANEXO N°/ANNEX NO: I

CERTIFICADO CE DE SISTEMA DE GARANTÍA DE CALIDAD TOTAL
de acuerdo con el Anexo IV (excepto punto 4) de la Directiva 98/79/CE

EC FULL QUALITY ASSURANCE SYSTEM CERTIFICATE
in accordance with Annex IV (except Section 4) of Directive 98/79/EC

Certificado n°/Certificate no	Fecha de validez/Date of validity		ON n°/NB no
2004 05 0442 CT	Desde/From	Hasta/To	0318
	20-05-2022	26-05-2025	

3.3 DIA.CHEMILUX Toxoplasma IgM

- RATOXOM.CE (100 tests)

3.4 DIA.CHEMILUX Toxoplasma IgG

- RATOXOG.CE (100 tests)

3.5 DIA.CHEMILUX Rubella IgM

- RARUBM.CE (100 tests)

3.6 DIA.CHEMILUX Rubella IgG

- RARUBG.CE (100 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 mayo de 2022

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS

 **agencia española de
medicamentos y
productos sanitarios**

Fdo. M^a Jesús Lamas Díaz

Agencia Española de Medicamentos y Productos Sanitarios (AEMPS)

Fecha de la firma: 19/05/2022

Puede comprobar la autenticidad del documento en la sede de la AEMPS: <https://localizador.aemps.es>

CSV: B 8 B Q W K 2 5 B 8



CORREO ELECTRÓNICO
on0318@aemps.es

Página 6 de 6

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel.: (+34) 91 822.57.87 / (+34) 91.822.59.97
Fax: (+34) 91.822.52.89

GMED certifie que le système de management de la qualité développé par
GMED certifies that the quality management system developed by

ELITECH CLINICAL SYSTEMS SAS
Zone Industrielle
61500 SEES FRANCE

pour les activités
for the activities

**Conception, production, contrôle et commercialisation de produits de chimie cliniques
pour le diagnostic in vitro. Validation de la combinaison réactifs et automates.
Distribution d'automates et de produits de chimie cliniques pour le diagnostic in vitro.**

*Design, production, control and sales of clinical chemistry products intended to be used
for in vitro diagnostics. Validation of the combination reagents and analyzers.
Distribution of clinical chemistry analyzers and products for in vitro diagnostics.*

réalisées sur le(s) site(s) de
performed on the location(s) of

ELITech Clinical Systems SAS
Zone industrielle - 61500 SEES - FRA

est conforme aux exigences des normes internationales
complies with the requirements of the international standards

NF EN ISO 13485 : 2016

Début de validité / Effective date : July 25th, 2023 (included)

Valable jusqu'au / Expiry date : July 27th, 2026 (included)

Etabli le / Issued on : July 25th, 2023



Accréditation n°4-0608
Liste des sites accrédités
et portée disponible sur
www.cofrac.fr

GMED N° 10462-8

Ce certificat est délivré selon les règles de certification GMED / This certificate is issued according to the rules of GMED certification

Renouvelle le certificat 10462-7



On behalf of the President
Marjorie PERRIMON
Certification Director



Medica Corporation
5 Oak Park Drive
Bedford, Massachusetts 01730
Tel 781 275 4892
Fax 781 275 2731
www.medicacorp.com

Declaration of Conformity

Product Name:

EasyLyte Analyzer and accessories per attachment

Model/Type:

Na/K, Na/K/Cl, Na/K/Li,
Na/K/Cl/Li, Na/K/Ca/pH,
Na/K/Cl/Ca/Li

Manufacturer

 Medica Corporation
5 Oak Park Drive, Bedford, Massachusetts, 01730, USA
Single Registration Number (SRN): US-MF-000037250

Representative

EC REP Emergo Europe, Prinsessegracht 20,
2514 AP The Hague, The Netherlands
Tel: +31 70 345 8570
Fax: +31 70 346 7299

Means of Conformity

Medica Corporation declares that the products listed are covered by Annex III of Directive 98/79/EC. These products are self-certified since they are for professional use only and are not listed on Annex II, List A or Annex II, List B of Directive 98/79/EC. In addition, they are in conformity with the Annex I, "Essential Requirements" and provisions of council Directive 98/79/EC for In Vitro Diagnostic Medical Devices, Directive 2011/65/EU Restriction of Hazardous Substance in Electrical and Electronic Equipment, and their corresponding amendments.

Place and Date: Bedford, Massachusetts, USA, 26 May 2022

Signature:



Name: Photios Makris, Ph.D.
Title: VP, Regulatory Affairs



Medica Corporation
5 Oak Park Drive
Bedford, Massachusetts 01730
Tel 781 275 4892
Fax 781 275 2731
www.medicacorp.com

Catalog No.	EasyLyte Analyzer and Accessories	EDMA Code	Class
2004	EasyLyte Analyzer, Na/K	21 07 11 02	General IVD device, not listed in IVD Annex II and not intended for self-testing
2014	EasyLyte Analyzer, Na/K/Cl	21 07 11 02	
2015	EasyLyte Analyzer, Na/K/Li	21 07 11 02	
2016	EasyLyte Analyzer, Na/K/Ca/pH	21 07 11 02	
2021	EasyLyte Analyzer, Na/K/Cl/Li	21 07 11 02	
2030	EasyLyte Analyzer, Na/K/Cl/Ca/Li	21 07 11 02	
C2004	EasyLyte Analyzer, Na/K	21 07 11 02	
C2014	EasyLyte Analyzer, Na/K/Cl	21 07 11 02	
C2015	EasyLyte Analyzer, Na/K/Li	21 07 11 02	
C2016	EasyLyte Analyzer, Na/K/Ca/pH	21 07 11 02	
C2030	EasyLyte Analyzer, Na/K/Cl/Ca/Li	21 07 11 02	
L2014	EasyLyte Analyzer, Na/K/Cl	21 07 11 02	
L2015	EasyLyte Analyzer, Na/K/Li	21 07 11 02	
L2016	EasyLyte Analyzer, Na/K/Ca/pH	21 07 11 02	
L2021	EasyLyte Analyzer, Na/K/Cl/Li	21 07 11 02	
2101	EasyLyte K ⁺ Electrode	11 04 01 06	
2102	EasyLyte Na ⁺ Electrode	11 04 01 07	
2103	EasyLyte Reference Electrode	11 04 04 01	
2113	EasyLyte Cl ⁻ Electrode	11 04 01 03	
2106	EasyLyte Lithium Electrode	11 04 01 04	
2150	EasyLyte Ca ⁺⁺ Electrode	11 04 01 02	
2151	EasyLyte pH Electrode	11 70 31 02	
2152	EasyLyte Disposable Reference Electrode	11 04 04 01	
2109	EasyLyte Solutions Pack, 400mL	11 04 04 02	
2120	EasyLyte Solutions Pack, 800mL	11 04 04 02	
2112	EasyLyte Plus Solutions Pack, 400mL	11 04 04 02	
2121	EasyLyte Plus Solutions Pack, 800mL	11 04 04 02	
2115	EasyLyte Lithium Solutions Pack, 400mL	11 04 04 02	
2122	EasyLyte Lithium Solutions Pack, 800mL	11 04 04 02	
2114	EasyLyte Calcium Solutions Pack, 400mL	11 04 04 02	
2123	EasyLyte Calcium Solutions Pack, 800mL	11 04 04 02	
2026	EasyLyte Na/K/Cl/Li Solutions Pack, 800mL	11 04 04 02	
2028	EasyLyte Na/K/Cl/Li Solutions Pack, 400mL	11 04 04 02	
2124	EasyLyte Na/K/Cl/Ca/Li Solutions Pack, 800mL	11 04 04 02	



Medica Corporation
5 Oak Park Drive
Bedford, Massachusetts 01730
Tel 781 275 4892
Fax 781 275 2731
www.medicacorp.com

Catalog No.	EasyLyte Analyzer and Accessories	EDMA Code	Class
2814	EasyQC Bi-Level Quality Control Kit	11 50 02 04	General IVD device, other than listed in IVD Annex II and other than intended for self-testing
2815	EasyQC Tri-Level Quality Control Kit	11 50 02 04	
L2026	EasyLyte Solutions Pack, Na/K/Cl/Li, 800mL	11 04 04 02	
L2112	EasyLyte Solutions Pack, Na/K/Cl, 400mL	11 04 04 02	
L2121	EasyLyte Solutions Pack, Na/K/Cl, 800mL	11 04 04 02	
L2122	EasyLyte Solutions, Na/K/Li Pack, 800mL	11 04 04 02	
L2123	EasyLyte Solutions Pack, Na/K/Ca/pH, 800mL	11 04 04 02	

EasyLyte EasyBloodGas EasyStat

Training Certificate

This is to certify that

Sorocovici Sergiu
Of Global Biomarketing Group
has completed training for the operation and service of the
EasyLyte, EasyBloodGas, and EasyStat analyzers.

November 25, 2004

Date



MEDICA

Randall Rollins

Signed: Randall Rollins
Technical Service Manager

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.com)

3) Product(s) (name, type or model/batch number, etc.):

Immunoassay products;

AccuBind® ELISA,

AccuLite® CLIA,

QSure® Control,

Instruments

see appendix

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

<u>Document No.</u>	<u>Title</u>
98/79/EC	<i>In vitro</i> Diagnostic Medical Devices Directive

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; 2021-09-20



(Place & date of issue (yyyy-mm-dd))

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

Appendix

Date: 2021-09-20

List of devices.

Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# QSure® Control	Item# Instru- ment	EDMS code	Risk Class	First date of CE-marking
Allergy & Anemia							
Ferritin Test System	2825-300A 2825-300B	2875-300A 2875-300B			12.07.01.02.00	Low	2005-11-11
Folate Test System	7525-300A 7525-300B	7575-300A 7575-300B			12.07.01.03.00	Low	2010-06-29
Immunoglobulin E (IgE) Test System	2525-300A 2525-300B	2575-300A 2575-300B			12.02.01.02.00	Low	2005-11-11
Transferrin Soluble Receptor (sTfR) Test System	8625-300A 8625-300B	8675-300A 8675-300B			12.07.01.06.00	Low	2010-06-29
Vitamin B-12 (Vit B12) Test System	7625-300A 7625-300B	7675-300A 7675-300B			12.07.02.04.00	Low	2011-09-26
Folate, Vitamin B-12 (Anemia Panel VAST) Test System	7825-300A 7825-300B	7875-300A 7875-300B			12.07.01.00.00	Low	2013-09-16
Autoimmune							
Anti-Cyclic Citrullinated Peptide IgG (Anti-CCP IgG) Test System	12725-300A 12725-300B	12775-300A 12775-300B			12.11.01.90.00	Low	2019-04-03
Anti-Thyroglobulin (Anti-Tg) Test System	1025-300A 1025-300B	1075-300A 1075-300B			12.10.03.04.00	Low	2005-11-11
Anti-Thyroperoxidase (Anti-TPO) Test System	1125-300A 1125-300B	1175-300A 1175-300B			12.10.03.01.00	Low	2005-11-11
Bone Metabolism & Growth							
Calcitonin Test System	9325-300A 9325-300B	9375-300A 9375-300B			12.06.03.02.00	Low	2019-04-03
Growth Hormone (hGH) Test System	1725-300A 1725-300B	1775-300A 1775-300B			12.06.04.02.00	Low	2005-11-11
Parathyroid Hormone (PTH) Test System	9025-300A 9025-300B	9075-300A 9075-300B			12.06.03.13.00	Low	2011-09-26
Parathyroid Hormone (PTH) 3rd & 2nd Gen (VAST) Test System	10025-300A 10025-300B	10075-300A 10075-300B			12.06.03.13.00	Low	2019-04-03
25(OH) Vitamin D Total Direct (Vit D-Direct) Test System	7725-300A 7725-300B	7775-300A 7775-300B			12.06.03.10.00	Low	2017-07-05
Cancer Markers							
Alpha-Fetoprotein (AFP) Test System	1925-300A 1925-300B	1975-300A 1975-300B			12.03.90.01.00	Low	2005-11-11
CA-125 Test System	3025-300A 3025-300B	3075-300A 3075-300B			12.03.01.06.00	Low	2005-11-11
CA 15-3 Test System	5625-300A 5625-300B	5675-300A 5675-300B			12.03.01.02.00	Low	2010-06-29
CA 19-9 Test System	3925-300A 3925-300B	3975-300A 3975-300B			12.03.01.03.00	Low	2005-11-11
Carcinoembryonic Antigen (CEA) Test System	1825-300A 1825-300B	1875-300A 1875-300B			12.03.01.31.00	Low	2005-11-11
Next Generation Carcinoembryonic Antigen	4625-300A	4675-300A			12.03.01.31.00	Low	2010-06-29

<i>Device types</i>	<i>Item# AccuBind® ELISA Microwells</i>	<i>Item# AccuLite® CLIA Microwells</i>	<i>Item# QSure® Control</i>	<i>Item# Instru- ment</i>	<i>EDMS code</i>	<i>Risk Class</i>	<i>First date of CE-marking</i>
(CEA-Next Gen) Test System	4625-300B	4675-300B					
Free β-Subunit Human Chorionic Gonadotropin (Free Beta hCG) Test System	2025-300A 2025-300B	2075-300A 2075-300B			12.03.01.90.00	Low	2005-11-11
Cardiac Markers							
CK-MB Test System	2925-300A 2925-300B	2975-300A 2975-300B			12.13.01.02.00	Low	2005-11-11
Digoxin (DIG) Test System	925-300A 925-300B	975-300A 975-300B			12.08.01.01.00	Low	2005-11-11
High Sensitivity CRP (hs-CRP) Test System	3125-300A 3125-300B	3175-300A 3175-300B			12.13.01.90.00	Low	2005-11-11
Myoglobin Test System	3225-300A 3225-300B	3275-300A 3275-300B			12.13.01.05.00	Low	2005-11-11
Troponin I (cTnI) Test System	3825-300A 3825-300B	3875-300A 3875-300B			12.13.01.07.00	Low	2005-11-11
Diabetes							
C-Peptide Test System	2725-300A 2725-300B	2775-300A 2775-300B			12.06.01.01.00	Low	2005-11-11
Insulin Test System	2425-300A 2425-300B	2475-300A 2475-300B			12.06.01.03.00	Low	2005-11-11
Rapid Insulin Test System	5825-300A 5825-300B				12.06.01.03.00	Low	2010-06-29
Insulin - C-Peptide (Diabetes Panel VAST)	7325-300A 7325-300B	7375-300A 7375-300B			12.06.01.03.00	Low	2005-11-11
Endocrine							
ACTH Test System	10625-300	10675-300			12.06.04.01.00	Low	2019-04-03
Aldosterone Test System	10125-300	10175-300			12.06.02.01.00	Low	2019-04-03
Leptin Test System	10925-300	10975-300			12.06.90.17.00	Low	2019-04-03
Fertility & Prenatal							
Anti-Müllerian Hormone (AMH) Test System	9725-300A 9725-300B	9775-300A 9775-300B			12.05.02.16.00	Low	2019-04-03
Folicle Stimulating Hormone (FSH) Test System	425-300A 425-300B	475-300A 475-300B			12.05.01.04.00	Low	2005-11-11
B-Human Chorionic Gonadotropin (hCG) Test System	825-300A 825-300B	875-300A 875-300B			12.05.02.05.00	Low	2005-11-11
B-Human Chorionic Gonadotropin Extended Range (hCG-XR) Test System	8825-300A 8825-300B	8875-300A 8875-300B			12.05.02.05.00	Low	2013-09-16
Rapid B-Human Chorionic Gonadotropin (Rapid-hCG) Test System	3325-300A 3325-300B				12.05.02.05.00	Low	2005-11-11
Inhibin A Test System	9525-300A 9525-300B	9575-300A 9575-300B			12.05.01.90.00	Low	2019-04-03
Inhibin B Test System	9625-300A 9625-300B	9675-300A 9675-300B			12.05.01.90.00	Low	2019-04-03
Luteinizing Hormone (LH) Test System	625-300A 625-300B	675-300A 675-300B			12.05.01.05.00	Low	2005-11-11
Pregnancy Associated Plasma Protein – A Mass Units (PAPP-A Mass Units) Test System	12625-300A 12625-300B	12675-300A 12675-300B			12.05.02.10.00	Low	2017-07-05
Prolactin Hormone (PRL) Test System	725-300A 725-300B	775-300A 775-300B			12.05.01.08.00	Low	2005-11-11

Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# QSure® Control	Item# Instru- ment	EDMS code	Risk Class	First date of CE-marking
Prolactin Hormone Sequential (PRLs) Test System	4425-300A 4425-300B	4475-300A 4475-300B			12.05.01.08.00	Low	2005-11-11
Human Chorionic Gonadotropin (hCG) , Human Prolactin (hPRL), Human Luteinizing Hormone (hLH), Follicle Stimulating Hormone (FSH) (Fertility Panel VAST) Test System	8325-300B 8325-300D 8325-300E	8375-300B 8375-300D 8375-300E			12.05.01.90.00	Low	2006-08-24
Alpha-Fetoprotein (AFP), Human Chorionic Gonadotropin (hCG), Unconjugated Estiol (u-E3) Triple Screen (Triple Screen Panel VAST) Test System	8525-300A 8525-300B	8575-300A 8575-300B			12.05.01.90.00	Low	2010-06-29
Infectious Diseases							
Anti-H. Pylori IgG (H. Pylori Ab IgG) Test System	1425-300A 1425-300B	1475-300A 1475-300B			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgM (H. Pylori Ab IgM) Test System	1525-300A 1525-300B	1575-300A 1575-300B			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgA (H. Pylori Ab IgA) Test System	1625-300A 1625-300B	1675-300A 1675-300B			15.01.04.03.00	Low	2005-11-11
Anti-SARS-CoV-2 (COVID-19) IgG Test System	11925-300A 11925-300B	11975-300A 11975-300B			15.04.80.90.00	Low	2020-08-25
Anti-SARS-CoV-2 (COVID-19) IgM Test System	11725-300A 11725-300B	11775-300A 11775-300B			15.04.80.90.00	Low	2020-08-25
Anti-SARS-CoV-2 (COVID-19) IgA Test System	11825-300A 11825-300B	11875-300A 11875-300B			15.04.80.90.00	Low	2020-08-25
Anti-SARS-CoV-2 (COVID-19) S1-RBD IgG Test System	12025-300A 12025-300B	12075-300A 12075-300B			15.04.80.90.00	Low	2021-09-20
D-Dimer Test System	9225-300A 9225-300B	9275-300A 9275-300B			13.02.05.03.00	Low	2020-08-25
Procalcitonin (PCT) Test System	1425-300A 1425-300B	1475-300A 1475-300B			12.06.90.16.00	Low	2017-07-05
Neonatal							
Neonatal 17OHP (N-17OHP) Test System	5525-300A 5525-300B				12.05.01.07.00	Low	2008-02-01
Neonatal (N-T4) Thyroxine Test System	2625-300A 2625-300B				12.04.01.12.00	Low	2005-11-11
Neonatal TBG (N-TBG) Test System	8925-300A 8925-300B				12.04.01.09.00	Low	2013-09-16
Neonatal TSH (N-TSH) Test System	3425-300A 3425-300B 3425-300D 3425-300E				12.04.01.90.00	Low	2005-11-11
Steroid							
Androstenedione (ANST) Test System	12425-300A 12425-300B	12475-300A 12475-300B			12.05.01.01.00	Low	2021-09-20
Cortisol Test System	3625-300A 3625-300B	3675-300A 3675-300B			12.06.02.04.00	Low	2005-11-11
Dehydroepiandrosterone (DHEA) Test System	7425-300A 7425-300B	7475-300A 7475-300B			12.05.01.02.00	Low	2011-09-26
Dehydroepiandrosterone Sulfate (DHEA-S) Test System	5125-300A 5125-300B	5175-300A 5175-300B			12.05.01.02.00	Low	2010-06-29
Estrone (E1) Test System	10325-300A 10325-300B	10375-300A 10375-300B			12.05.02.04.00	Low	2019-04-03

<i>Device types</i>	<i>Item# AccuBind® ELISA Microwells</i>	<i>Item# AccuLite® CLIA Microwells</i>	<i>Item# QSure® Control</i>	<i>Item# Instru- ment</i>	<i>EDMS code</i>	<i>Risk Class</i>	<i>First date of CE-marking</i>
Estradiol (E2) Test System	4925-300A 4925-300B	4975-300A 4975-300B			12.05.01.03.00	Low	2010-06-29
Unconjugated Estiol (u-E3) Test System	5025-300A 5025-300B	5075-300A 5075-300B			12.05.02.02.00	Low	2010-06-29
Progesterone Test System	4825-300A 4825-300B	4875-300A 4875-300B			12.05.01.06.00	Low	2010-06-29
17-OH Progesterone (17-OHP) Test System	5225-300A 5225-300B	5275-300A 5275-300B			12.05.01.07.00	Low	2010-06-29
17-OH Progesterone SI (17-OHP-SI) Test System	9925-300A 9925-300B	9975-300A 9975-300B			12.05.01.07.00	Low	2010-10-18
Sex Hormone Binding Globulin (SHBG) Test System	9125-300A 9125-300B	9175-300A 9175-300B			12.05.01.09.00	Low	2013-09-16
Testosterone Test System	3725-300A 3725-300B	3775-300A 3775-300B			12.05.01.10.00	Low	2007-11-01
Free Testosterone Test System	5325-300A 5325-300B	5375-300A 5375-300B			12.05.01.10.00	Low	2010-06-29
Thyroid							
Total Triiodothyronine (tT3) Test System	125-300A 125-300B 125-300D 125-300E	175-300A 175-300B 175-300D 175-300E			12.04.01.05.00	Low	2005-11-11
Free Triiodothyronine (fT3) Test System	1325-300A 1325-300B 1325-300A 1325-300B	1375-300A 1375-300B 1375-300D 1375-300E			12.04.01.01.00	Low	2005-11-11
Total Triiodothyronine (tT3 SBS) Test System	8125-300A 8125-300B	8175-300A 8175-300B			12.04.01.01.00	Low	2010-06-29
Rapid Total Triiodothyronine (Rapid -tT3) Test System	11225-300A 11225-300B				12.04.01.01.00	Low	2017-07-05
T3-Uptake (T3U) Test System	525-300A 525-300B	575-300A 575-300B			12.04.01.06.00	Low	2005-11-11
Thyroxine (tT4) Test System	225-300A 225-300B 225-300D 225-300E	275-300A 275-300B 275-300D 275-300E			12.04.01.07.00	Low	2005-11-11
Free Thyroxine (fT4) Test System	1225-300A 1225-300B 1225-300D 1225-300E	1275-300A 1275-300B 1275-300D 1275-300E			12.04.01.02.00	Low	2005-11-11
Total Thyroxine (tT4 SBS) Test System	8225-300A 8225-300B	8275-300A 8275-300B			12.04.01.01.00	Low	2010-06-29
Rapid Total Thyroxine (Rapid -tT4) Test System	11125-300A 11125-300B				12.04.01.01.00	Low	2017-07-05
Thyrotropin (TSH) Test System	325-300A 325-300B 325-300D 325-300E	375-300A 375-300B 375-300D 375-300E			12.04.01.11.00	Low	2005-11-11
Rapid TSH Test System	6025-300A 6025-300B	6075-300A 6075-300B			12.04.01.11.00	Low	2010-06-29
Thyroxine-Binding Globulin (TBG) Test System	3525-300A 3525-300B	3575-300A 3575-300B			12.04.01.09.00	Low	2005-11-11
Thyroglobulin (Tg) Test System	2225-300A	2275-300A			12.04.01.08.00	Low	2005-11-11

Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# QSure® Control	Item# Instru- ment	EDMS code	Risk Class	First date of CE-marking
	2225-300B	2275-300B					
Total Thyroxine (tT4), Total Triiodothyronine (tT3) & Thyroid Stimulating Hormone (TSH) (Thyroid Panel VAST) Test System	8025-300B 8025-300D 8025-300E	8075-300B 8075-300D 8075-300E			12.04.01.01.00	Low	2005-11-11
Free Thyroxine (fT4), Free Triiodothyronine (fT3) & Thyroid Stimulating Hormone (TSH) (Free Thyroid Panel VAST) Test System	7025-300B 7025-300D 7025-300E	7075-300B 7075-300D 7075-300E			12.04.01.01.00	Low	2010-06-29

Miscellaneous Controls							
Anti-H. Pylori Control (IgA, IgG, IgM) – Positive & Negative			HPC-300		12.50.01.16.00	Low	2013-09-16
Anti-Tg & Anti-TPO Control – Positive & Negative			AIT-101		12.50.01.16.00	Low	2010-06-29
Maternal Control – (AFP, uE3, hCG, Free beta hCG) Tri Level			MC-300		12.50.01.16.00	Low	2010-06-29
TBG Control – Tri-Level			TBG-300		12.50.01.16.00	Low	2013-09-16
Tg Control – Tri-Level			TG-300		12.50.01.16.00	Low	2010-06-29
Tumor Marker Control – (CA 125, CA 15-3, CA 19-9) Tri-Level			TMC-300		12.50.01.16.00	Low	2013-09-16

Miscellaneous Instruments							
Autoplex® ELISA & CLIA Analyzer				IN006	21.02.10.01	Low	2010-06-29
Autoplex® G2 ELISA & CLIA Analyzer				IN006-2	21.02.10.01	Low	2013-09-16
Autoplex® G3 ELISA & CLIA Analyzer				IN006-3	21.02.10.01	Low	2017-07-05
NeoEldex® ELISA Analyzer				IN009	21.02.10.01	Low	2011-09-26
Impulse® 3 CLIA Analyzer				IN007	21.02.10.01	Low	2010-06-29
NeoLumax® CLIA Analyzer				IN010	21.02.10.01	Low	2011-09-26
LuMatic® CLIA Analyzer				IN008	21.02.10.01	Low	2011-09-26
PrisMatic® ELISA Analyzer				IN013	21.02.10.01	Low	2013-09-16
PlateWash - Immunoassay Washer				IN002	21.02.10.01	Low	2010-06-29
TITIN® ELISA & CLIA Analyzer				IN015-EC	21.02.10.01	Low	2017-07-05
TITIN® ELISA Analyzer				IN015-E	21.02.10.01	Low	2017-07-05
TITIN-s® ELISA & CLIA Analyzer				IN016-EC	21.02.10.01	Low	2017-07-05
TITIN-s® ELISA Analyzer				IN016-E	21.02.10.01	Low	2017-07-05



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗДРАВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 22 января 2016 года № ФСР 2012/14183

На медицинское изделие

**Набор реагентов для клинического анализа спинномозговой жидкости
(«ДИАХИМ-ЛИКВОР») по ТУ 9398-056-27428909-2012**

Настоящее регистрационное удостоверение выдано

**Общество с ограниченной ответственностью "Научно-производственная фирма
"АБРИС+" (ООО "НПФ" АБРИС+), Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж**

Производитель

**Общество с ограниченной ответственностью "Научно-производственная фирма
"АБРИС+" (ООО "НПФ" АБРИС+), Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж**

Место производства медицинского изделия

192019, Санкт-Петербург, ул. Профессора Качалова, д. 15а, лит. А

Номер регистрационного досье № РД-9561/60865 от 14.12.2015

Вид медицинского изделия -

Класс потенциального риска применения медицинского изделия 1

Код Общероссийского классификатора продукции для медицинского изделия 93 9816

Настоящее регистрационное удостоверение имеет приложение на 1 листе

приказом Росздравнадзора от 22 января 2016 года № 341
допущено к обращению на территории Российской Федерации.

**Руководитель Федеральной службы
по надзору в сфере здравоохранения**



М.А. Мурашко

0017000

**ПРИЛОЖЕНИЕ
К РЕГИСТРАЦИОННОМУ УДОСТОВЕРЕНИЮ
НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ**

от 22 января 2016 года № ФСР 2012/14183

Лист 1

На медицинское изделие

**Набор реагентов для клинического анализа спинномозговой жидкости
(«ДИАХИМ-ЛИКВОР») по ТУ 9398-056-27428909-2012:**

- Реактив Самсона - готов к применению - 1 флакон 10 мл.
- Карболовая кислота - готов к применению - 1 флакон 2,5 г.
- Аммоний сернокислый - готов к применению - 1 флакон (пакет) 85 г.

З

Руководитель Федеральной службы
по надзору в сфере здравоохранения



М.А. Мурашко

0014999



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗДРАВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 22 января 2016 года № ФСР 2010/07198

На медицинское изделие

Набор реагентов для клинического анализа кала («Диахим-Набор реагентов для клинического анализа кала») по ТУ 9398-033-27428909-2009

Настоящее регистрационное удостоверение выдано

Общество с ограниченной ответственностью "Научно-производственная фирма "АБРИС+" (ООО "НПФ"АБРИС+"), Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж

Производитель

Общество с ограниченной ответственностью "Научно-производственная фирма "АБРИС+" (ООО "НПФ"АБРИС+"), Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж

Место производства медицинского изделия

192019, Санкт-Петербург, ул. Профессора Качалова, д. 15а, лит. А

Номер регистрационного досье № РД-9543/60775 от 14.12.2015

Вид медицинского изделия -

Классе потенциального риска применения медицинского изделия 2а

Код Общероссийского классификатора продукции для медицинского изделия 93 9816

приказом Росздравнадзора от 22 января 2016 года № 354
допущено к обращению на территории Российской Федерации.

Руководитель Федеральной службы
по надзору в сфере здравоохранения

М.А. Мурашко

0017228



Санкт-Петербург



ВСЕРОССИЙСКИЙ НАУЧНО-ИССЛЕДОВАТЕЛЬСКИЙ
ИНСТИТУТ СЕРТИФИКАЦИИ
ОТКРЫТОЕ АКЦИОНЕРНОЕ ОБЩЕСТВО
(ОАО "ВНИИС")

Электрический пер., д.3/10, строение 1,
г. Москва, 125057

Телефон: (499) 253 70 06 Факс: (499) 253 33 50
<http://www.vniis.ru> E-mail: vniis@vniis.ru

Исх. № 102-К/1010 от 20.11.15

Генеральному директору
ООО «НПФ «АБРИС+»
В.М. Марковскому
196084, г. Санкт-Петербург,
ул. Цветочная, д.16, БЦ «Осиповф»,
офис 207
тел.: (812) 458-44-07

На № 166
от 10.11.2015г.

На Ваш запрос о принадлежности к объектам обязательного подтверждения соответствия наборов реагентов для клинической лабораторной диагностики и иммунохимических анализов, согласно приложению, сообщаем следующее.

Продукция, указанная в приложении, может быть отнесена по Общероссийскому классификатору продукции ОК 005-93 к позициям: «Наборы реагентов для клинической лабораторной диагностики» (код ОКП 93 9816), «Наборы реагентов для радиоиммунологического и других видов иммунохимических анализов» (код ОКП 93 9817).

Продукция, указанная в приложении к настоящей справке, не включена в «Единый перечень продукции, подлежащей обязательной сертификации» и «Единый перечень продукции, подтверждение соответствия которой осуществляется в форме принятия деклараций о соответствии», утвержденные Постановлением Правительства Российской Федерации от 01.12.09 г. № 982 (с изменениями), а также не включена в «Единый перечень продукции, подлежащей обязательной оценке (подтверждению) соответствия в рамках Таможенного союза с выдачей единых документов», утвержденный Решением Комиссии Таможенного союза от 07.04.2011г. № 620 (с изменениями), и для нее не требуется представление сертификата соответствия или декларации о соответствии.

Одновременно сообщаем, что продукция, указанная в приложении не подпадает под действие вступивших в силу технических регламентов Таможенного союза, и для нее не требуется представление документов о подтверждении соответствия требованиям этих технических регламентов.

Настоящая справка действительна до внесения изменений в документы, устанавливающие необходимость проведения обязательного подтверждения соответствия данной продукции.

Приложение: на 7 л. в 1 экз.

Руководитель научного направления

Круглосуточный автоинформатор: (499) 253 00 78
телефоны для справок: (499) 253 03 68, (499) 253 03 79
факсы: (499) 253 00 85, (499) 253 68 55



Система качества ВНИИС сертифицирована



54.	Раствор бриллиантового крейлового синего для окраски ретикулоцитов (Диахим-Гемистейн-РПЦ)	
55.	Фиксатор-краситель эозин метиленовый синий по Май-Грюнвальду (Диахим-Гемистейн-М-Г)	
56.	Фиксатор-краситель эозин метиленовый синий типа Лейшмана (Диахим-Гемистейн-Л)	
57.	Краситель азур-эозин по Романовскому (Диахим-Гемистейн-Р)	
58.	Набор реагентов для окраски по Граму «Диахим-набор для окраски по Граму»	
59.	Набор реагентов для окраски по Циль-Нильсену «Диахим-набор для окраски по Циль-Нильсену»	
60.	Набор реагентов для клинического анализа кала ("Диахим-Набор реагентов для клинического анализа кала")	
61.	Набор реагентов для исследования фекалий по методу Като ("Диахим-КАТО")	
62.	Набор реагентов для исследования на гельминты по Рабиновичу ("ДИАХИМ-Набор реагентов для исследования на гельминты по Рабиновичу")	- Клеевой состав - готов к применению - 1 флакон 50 мл. - Стекланные палочки (лопатки) - 50 шт.
63.	Набор реагентов для окраски урогенитальных мазков по Папаниколау ("ДИАХИМ-ПАП")	- Гематоксилин Гарриса - 1 фл (100 мл). - Краситель Папаниколау - 1 фл (100 мл). - Краситель оранжевый G6 - 1 фл (100 мл). - Литий углекислый - 1 фл (3 мл).
64.	Набор реагентов для быстрого дифференцированного окрашивания биопрепаратов ("ДИАХИМ-ДИФФ-КВИК")	- Раствор № 1 (фиксатор) - 100 мл - Раствор № 2 («розовый») - 100 мл - Раствор № 3 («синий») - 100 мл - Буферная смесь - 1 флакон
65.	Краситель для цитологических исследований (Гематоксилин Майера) «ДИАХИМ-ЦИТОСТЕЙН-ГМ»	- флакон с красителем; - водный раствор гематоксилина; - водный раствор алюмокалиевых квасцов; - водный раствор йоднокислого калия; - глицерин.
66.	Краситель для цитологических исследований (Гематоксилин Караши) «ДИАХИМ-ЦИТОСТЕЙН-ГК»	- флакон с красителем; - водный раствор гематоксилина; - водный раствор алюмокалиевых квасцов; - водный раствор йоднокислого калия; - хлоралгидрат.

Руководитель научного направления ВНИИС





ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗДРАВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 31 декабря 2015 года № ФСР 2010/08734

На медицинское изделие

**Набор реагентов для исследования фекалий по методу Като («Диахим-КАТО»)
по ТУ 9398-042-27428909-2010**

Настоящее регистрационное удостоверение выдано

**Общество с ограниченной ответственностью "Научно-производственная фирма
"АБРИС+" (ООО "НПФ" АБРИС+)", Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж**

Производитель

**Общество с ограниченной ответственностью "Научно-производственная фирма
"АБРИС+" (ООО "НПФ" АБРИС+)", Россия,
196084, Санкт-Петербург, ул. Цветочная, д. 16, лит. М, 2-й этаж**

Место производства медицинского изделия

192019, Санкт-Петербург, ул. Профессора Качалова, д. 15а, лит. А

Номер регистрационного досье № РД-9559/60784 от 14.12.2015

Вид медицинского изделия 139290

Класс потенциального риска применения медицинского изделия 2а

Код Общероссийского классификатора продукции для медицинского изделия 93 9816

приказом Росздравнадзора от 31 декабря 2015 года № 10009
допущено к обращению на территории Российской Федерации.

**Руководитель Федеральной службы
по надзору в сфере здравоохранения**



М.А. Мурашко

0016928