



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	CMV IgG CODE: CMVG.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY	AEMPS – n° 0318
(EC) CERTIFICATE(S)	<ul style="list-style-type: none"><li>FULL QUALITY ASSURANCE SYSTEM N° 2004 05 0442 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318</li><li>UNI EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318</li></ul>

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – MAY 2018
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018



Dia.Pro  
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# EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	Chlamydia Trachomatis IgG CODE: CTG.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY	AEMPS – n° 0318
(EC) CERTIFICATE(S)	<ul style="list-style-type: none"><li>FULL QUALITY ASSURANCE SYSTEM N° 2004 05 0442 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318</li><li>UNI EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318</li></ul>

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2009
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – MAY 2018
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018

Deltalab, S.L. defines and makes public its commitment with the Standard ISO 9001:2015 Quality Management Systems, ISO 14001:2015 Environmental Management Systems and ISO 13485:2016 Medical devices – Quality Management Systems, with the aim to create value and satisfy all its interested parties:

- Shareholders
- Members of the organisation
- Customers and suppliers
- All members of the surrounding community

The development of this Integrated Management System Policy is carried out with the philosophy of Continuous Improvement and with the support of all the processes described in our Integrated Management System, in order to achieve the following objectives:

1. Become leaders in the design and manufacture of single use products for the laboratory.
2. Bring solutions to cover the current and future customer needs, related to:
  - Design, manufacture and sale of laboratory material for the collection, transport and conservation of samples for microbiology, molecular biology, haematology, biochemistry, histology, microscopy and colorimetric analysis.
  - Design, manufacture and sale of sterile and non-sterile medical devices for the collection, transport and conservation of biological samples for clinical and IVD analysis.
  - Commercialization of diagnosis reagents, equipment and instrumentation for laboratory and equipment for the storage of prepared samples, cryogenic stored samples, general labware and industrial packages.
  - Commercialization of personal care, cosmetics and dietetic products
3. Maintain a constant growth, both in local and international markets, by means of mergers, acquisitions and by launching new products.
4. Achieve the full satisfaction of our customers, by means of a strict compliance to the agreements and expectations agreed with them, as well as the excellence in the service.
5. Reach a high level of innovation of our products and processes, in cooperation with universities, research centers, key opinion leaders and experts, both local and international.
6. Fulfil the legislation and regulatory requirements applicable to the activities carried out by the company, including those applicable to the quality of products and the environmental management.
7. Commit ourselves with the environmental protection, including the prevention of pollution.
8. Achieve and keep a high motivation and involvement of all members of the organisation, suppliers, distributors and customers, by fulfilling the highest Quality and environmental protection standards.
9. Improve the working conditions of all employees and ensure the technical capacity of the personnel by giving them the adequate training with the aim to achieve the required competence.
10. Establish a close relationship with the suppliers and guarantee the maximum quality of materials supplied by means of quality agreements.

The Integrated Management System is periodically reviewed to define the required actions to ensure that:

- ✓ The System is efficient, so that it is a tool for the routine of all the members of the organisation.
- ✓ The customer needs and requirements are duly identified, and their expectations are always met.
- ✓ All members of the organisation are familiar with and know the objectives and policy of the Integrated Management System, and that adequate training plans are defined to achieve them.
- ✓ Encourage the Continuous Improvement Philosophy, both related to Quality and Environmental Management.

This Policy is made available for the public and all interested parties.

JOSEP SAEZ  
Managing Director  
January 2019



Certificate ES16/20725

#### The management system of

**DELTALAB GROUP**  
**DELTALAB, S.L., KEYLAB, S.L.U.,**  
**NIRCO, S.L., ENVASES FARMACÉUTICOS, S.A.**

Pol. Ind. La Llana  
Plaza de la Verneda, 1  
08191 Rubí, Barcelona

has been assessed and certified as meeting the requirements of



ISO 9001:2015

For the following activities

Design, manufacture and sale of laboratory material for the collection, transport and conservation of samples for microbiological, molecular biology, haematology, biochemistry, histology, microscopy and colorimetric analysis, general labware, containers and healthcare products. Manufacture and commercialization of consumables for the laboratory. Commercialization and distribution of equipment for the storage of prepared samples, cryogenic stored samples, syringes, general labware and industrial packages. Commercialization and distribution of equipment and instrumentation for the laboratory, diagnostic kits, healthcare products, cosmetics and food for special medical purposes. Commercialization, distribution, installation and technical service of equipment and instrumentation for the laboratory.

This certificate is valid from  
11 October 2019 until 11 October 2022.  
Issue 4. Company certified since October 2010.  
Certified with SGS since 11 October 2016

This is a multisite certification. See following page(s).

Authorised by

*[Signature]*

## Certification Management



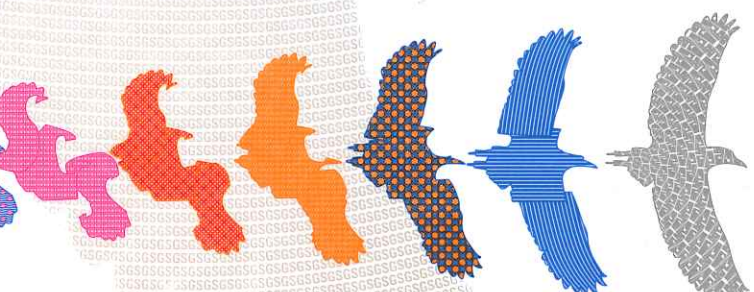
 ENAC  
CERTIFICACIÓN  
Nº 05/C-SC001

**SGS INTERNATIONAL CERTIFICATION SERVICES IBERICA, S.A.U.**  
C/Trespuentes, 29 28042 Madrid España  
t 3491 313 8115 f 34 91 313 8102 [www.sgs.com](http://www.sgs.com)

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S G S S G S & G





Certificate ES16/20725

**DELTALAB GROUP**  
**DELTALAB, S.L., KEYLAB, S.L.U.,**  
**NIRCO, S.L., ENVASES FARMACÉUTICOS, S.A.**

**ISO 9001:2015**

Issue 4



Sites where these activities are totally or partially carried out

**DELTALAB, S.L.**

Pol. Ind. La Llana, Plaza de la Verneda, 1 – 08191 Rubí, Barcelona (España)

Design, manufacture and sale of laboratory material for the collection, transport and conservation of samples for microbiological, molecular biology, haematology, biochemistry, histology, microscopy and colorimetric analysis. Commercialization of equipment for the storage of prepared samples, cryogenic stored samples, general labware and industrial packages.

Commercialization of equipment and instrumentation for the laboratory, diagnostic kits, healthcare products, cosmetics and food for special medical purposes.

**KEYLAB, S.L.U.**

Pol. Ind. La Llana, Avda. de la Llana, 115-117 – 08191 Rubí -Barcelona (España)

Design, manufacture and sale of laboratory material for the collection, transport and conservation of samples for microbiological, molecular biology, haematology, biochemistry, histology, microscopy and colorimetric analysis. Commercialization of equipment for the storage of prepared samples, cryogenic stored samples, general labware and industrial packages.

Commercialization of equipment and instrumentation for the laboratory, diagnostic kits, healthcare products, cosmetics and food for special medical purposes.



**NIRCO, S.L.**

Pol. Ind. Expansión, Puerto de Navafría, 12 - 28935 Móstoles -Madrid (España)

Pol. Ind. La Llana, Avda. de la Llana, 115-117 – 08191 Rubí -Barcelona (España)

Manufacture and commercialization of consumables for the laboratory.  
Commercialization and distribution of diagnostic kits

Commercialization, distribution, installation and technical service of equipment and instrumentation for the laboratory.

**ENVASES FARMACÉUTICOS, S.A.**

C/ Paralela, 15 - 28860 Paracuellos de Jarama (Madrid)

Design, manufacture and commercialization of laboratory material for the collection, transport and conservation of samples for analysis, laboratory material for general use, containers and products for personal care

Commercialisation and distribution of laboratory material for general use, products and equipment for personal care, syringes and cosmetic products.

## Certificate ES16/20725.01

 SGS

# **DELTALAB, S.L.**

Pol. Ind. La Llana  
Plaza de la Verneda, 1  
08191 Rubí, Barcelona

has been assessed as part of the management system of DELTALAB GROUP certified organization as meeting the requirements of

# ISO 9001:2015

For the following activities



**Design, manufacture and sale of laboratory material for the collection, transport and conservation of samples for microbiological, molecular biology, hematology, biochemistry, histology, microscopy and colorimetric analysis.**

**Commercialization of equipment for the storage of prepared samples, cryogenic stored samples, general labware and industrial packages.**

**Commercialization of equipment and instrumentation for the laboratory, diagnostic kits, healthcare products, cosmetics and food for special medical purposes.**

in / from the following sites

Pol. Ind. La Llana, Plaza de la Verneda, 1 - 08191 Rubí (Barcelona)

Valid from

September 2022.

This document is part of Certificate ES16/20725.  
The validity of this document is subject to the certificate.



 ENAC  
CERTIFICACIÓN  
Nº 05 / C - SC001

Authorized by

  
Dr. S. S.  
Sankararaman

## Certification Management

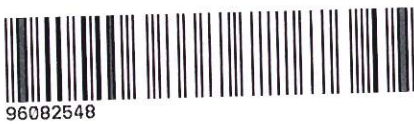
**SGS INTERNATIONAL CERTIFICATION SERVICES IBERICA, S.A.U.**  
C/Trespuentes, 29. 28042 Madrid. España.  
t 34 91 313 8115 f 34 91 313 8102 [www.sgs.com](http://www.sgs.com)

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S G S S G S C S





# Ministero della Salute

DIPARTIMENTO DELLA PROGRAMMAZIONE E DELL'ORDINAMENTO DEL SERVIZIO  
SANITARIO NAZIONALE

DIREZIONE GENERALE DEI DISPOSITIVI MEDICI, DEL SERVIZIO FARMACEUTICO  
E DELLA SICUREZZA DELLE CURE  
UFFICIO IV ex DGFDL – DIAGNOSTICI IN VITRO

I.5.l.e.2/IV/2011/37

VISTA la direttiva 98/79/CE relativa ai dispositivi medico-diagnostici in vitro;

VISTO il D.lgs. n .332/2000 recante attuazione della direttiva 98/79/CE;

VISTA l'istanza del 29/09/2011 presentata dalla ditta Dia.Pro Diagnostic Biopros Srl con sede in Via G.Carducci, 27 – 20099 Sesto San Giovanni (MI) – C.F./P.Iva 11924660159;

CONSIDERATO che la ditta istante ha effettuato i versamenti richiesti dal D.M. 24 Maggio 2004;

VISTI gli atti d'ufficio;

HAVING REGARD to 98/79/EC directive concerning the in vitro diagnostic medical-devices;

HAVING REGARD to legislative Decree (D.lgs.)n. 332/2000 reporting the accomplishment of 98/79/EC Directive;

HAVING REGARD to the request dated 29/09/2011 submitted by the company Dia.Pro Diagnostic Biopros Srl con with legal site in Via Columella, 31 – 20128 Milano – C.F. and P.Iva 11924660159;

WHEREAS this company paid the fees required by Ministerial Decree (D.M.) May 24, 2004;  
HAVING REGARD to the official deeds;

## SI ATTESTA IT IS ATTESTED

che la ditta, Dia.Pro Diagnostic Biopros Srl con sede in Via G.Carducci, 27 – 20099 Sesto San Giovanni (MI) – C.F./P.Iva 11924660159, ha prodotto e marcato CE, come dispositivo medico- diagnostico in vitro, secondo le procedure previste dalla direttiva 98/79/CE, il prodotto:

that the Company Dia.Pro Diagnostic Biopros Srl located in Via G.Carducci, 27 – 20099 Sesto San Giovanni (MI) – C.F./P.Iva 11924660159, manufactured and affixed CE marking as in vitro diagnostic medical device, according to the Directive 98/79/EC, the following product:

### DP-9 DIA.BLOOD INSTRUMENT

Il suddetto prodotto, in base all'art. 4 della direttiva 98/79/CE, è di libera circolazione e può essere messo in commercio in Italia e in tutto il territorio dell'Unione Europea.



Si rilascia il presente attestato su richiesta dell'interessato per gli usi consentiti dalla legge e per l'esportazione nei paesi extra UE.

*The above mentioned product, according to the art. 4 of 98/79/EC directive, can freely circulate and can be commercialized in Italy and in the whole of the European Union. This certificate is issued on the interested company's request according to the law and to export to non-European countries*



IC/CM

**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/EC***

**PRÓRROGA/EXTENSION — Fecha inicial/ Initial date: 15/03/2004**

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

<b>Certificado nº/Certificate no</b>	<b>Fecha de validez/Date of validity</b>	<b>ON nº/NB no</b>
<b>2004 03 0425 ED</b>	<b>Desde/From 26/11/2018 Hasta/To 18/11/2023</b>	<b>0318</b>

**A favor de /In favour of:**

**Fabricante/Manufacturer:**

**Nombre/Name:** DIA. Pro Diagnostic Bioprobe 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 Bioprobe 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, 23 de noviembre de 2018

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



agencia española de  
medicamentos y  
productos sanitarios

Fdo. Mª Jesús Lamas Díaz

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

Localizador: 3P6PS5XA6C

Puede comprobar la autenticidad del documento en la aplicación Localizador de la Web de la AEMPS  
**CORREO ELECTRÓNICO**  
on0318@aemps.es

Página 1 de 2

**ORGANISMO NOTIFICADO 0318**

C/ CAMPEZO, 1 - EDIFICIO 8

28022 MADRID

Tel.: (+34) 902.101.322 /(+34) 91.822.59.97

Fax: (+34) 91.822.52.89



**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/EC**  
**PRÓRROGA/EXTENSION — Fecha inicial/ Initial date: 15/03/2004**  
**Fecha de última prórroga/ Last extension date: 27/11/2013**

Certificado nº/Certificate no	Fecha de validez/Date of validity	ON nº/NB no
<b>2004 03 0425 ED</b>	Desde/From 26/11/2018 Hasta/To 18/11/2023	<b>0318</b>

A favor de/*In favour of:*

**Fabricante/Manufacturer:**

Nombre/*Name:* Dia. Pro Diagnostic Bioproses 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]

**HBe Ag & Ab ELISA cualitativo / ELISA qualitative**

- HBE.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

 agencia española de medicamentos y productos sanitarios

Fdo. M<sup>a</sup> Jesús Lamas Díaz

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

Localizador: 3P6PS5XA6C

Puede comprobar la autenticidad del documento en la aplicación Localizador de la Web de la AEMPS

**CORREO ELECTRÓNICO**

on0318@aemps.es

Página 2 de 2

**ORGANISMO NOTIFICADO 0318**

C/ CAMPEZO, 1 - EDIFICIO 8

28022 MADRID

Tel.: (+34) 902.101.322 /(+34) 91.822.59.97

Fax: (+34) 91.822.52.89



Dia.Pro  
Diagnostic  
BioProbes

# EC DECLARATION OF CONFORMITY

<b>MANUFACTURER</b>	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
<b>PRODUCT</b>	<b>HSV1&amp;2 IgG</b> CODE: <b>HSVG.CE</b> (96 tests)
<b>CLASSIFICATION</b>	GENERAL IVD
<b>CONFORMITY ASSESSMENT ROUTE</b>	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.**

<b>ISO CERTIFICATE</b>	UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY AEMPS (AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS)
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<b>PLACE &amp; DATE OF FIRST ISSUE</b>	MILANO – MARCH 2004
<b>PLACE &amp; DATE OF CURRENT ISSUE</b>	SESTO SAN GIOVANNI (MI) – MARCH 2019
<b>SIGNATURE</b> <b>Legal Representative</b> <b>Dr.ssa Fiorenza Scozzesi</b>	

Rev: 05/2018

**LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS**  
**THE AGENCY ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS**

otorga el certificado número

grants the certificate no.

**2013 11 0039 EN**

según la norma

in accordance with the standard

**UNE-EN ISO 13485:2018**

**(EN ISO 13485: 2016 & ISO 13485: 2016)**

**Productos Sanitarios: Sistemas de Gestión de Calidad – Requisitos para fines reglamentarios**

*Medical devices – Quality management systems - Requirements for regulatory purposes*

a la empresa / to the company

**Dia.Pro Diagnostic Bioprobe S.r.l.**

Sede social y de fabricación/ Headquarters and manufacturing facility

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

Para las siguientes actividades / For the following activities:

**Diseño, desarrollo y producción de reactivos y productos reactivos, calibradores y materiales de control para inmunoquímica, microbiología, inmunología infecciosa y técnicas de biología molecular.**

**Diseño, desarrollo, producción y servicio técnico de instrumentos y software para diagnóstico *in vitro*.**

*Design, development and manufacturing of reagents, reagent products, calibrators and control materials for immunochemistry, microbiology, infectious immunology and molecular biology techniques.*

*Design and development, management of production and technical servicing of instruments and software for "in vitro" diagnostic.*

**Modificaciones de alcance: Ver Anexo I / see Annex I**

**Fecha de validez/ Date of validity: Desde/ From: 8-03-2019 Hasta/To: 17-12-2021**

**Certificación inicial/ Initial certification date: 27-11-2013**

**Renovación / Renewal of certification date: 8-03-2019**

Madrid, 08 de marzo de 2019

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



Fdo. M<sup>a</sup> Jesús Lamas Díaz



## ANEXO I / ANNEX I

### CERTIFICADO UNE-EN ISO 13485:2018/ UNE-EN ISO 13485:2018 CERTIFICATE

#### Modificaciones del alcance / Scope modifications:

Fecha/Date	Descripción de la modificación/ Modification description
<b>18-12-2018</b>	<p>Cambio en la descripción del tipo de técnica en el ámbito tecnológico (inmunología infecciosa y técnicas de biología molecular). Cambio del nivel de detalle en la descripción del ámbito tecnológico</p> <p><i>Change in the description of the method of analysis in the technological scope (infectious immunology and molecular biology techniques). Change in the level of detail of the technological scope description.</i></p>
<b>8-03-2019</b>	<p>Ampliación del ámbito tecnológico para incluir:            Inmunoquímica y microbiología            Instrumentos y software para diagnóstico "in vitro".</p> <p>Modificación del alcance para incluir la actividad de asistencia técnica para Instrumentos y software para diagnóstico "in vitro".</p> <p><i>Extension of technological scope:            Immunochemistry and Microbiology            Instruments and software for "in vitro" diagnostic            Modification of the scope to include the activity of technical servicing of instruments and software for "in vitro" diagnostic</i></p>

Madrid, 08 de marzo de 2019

DIRECTORA DE LA AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS



Fdo. Mª Jesús Lamas Díaz



## **DECLARATION DE CONFORMITE CE**

Nous, ELITech Clinical Systems SAS, zone industrielle 61500 SEES France, déclarons sous notre seule responsabilité que les réactifs référencés dans la liste ci-jointe (2 pages), 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.

Ces dispositifs sont classés dans la catégorie « autre dispositif » puisqu'ils n'appartiennent ni à la liste A et liste B de l'annexe II et ni à la classe des autotests.

Cette déclaration est basée sur le contenu de chaque dossier technique et s'appuie sur la certification de notre système qualité selon la norme NF EN ISO 13485 : 2016 (Certification valable jusqu'au 27 juillet 2023).

## **DECLARATION OF EC CONFORMITY**

*We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents such as listed attached (2 pages), 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.*

*These devices are classified in the "other device" category since they do not belong neither to list A or list B of annex II nor to self-testing class.*

*This declaration is based on the contents of each 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 27<sup>th</sup>, 2023).*

## **DECLARACIÓN CE DE CONFORMIDAD**

*Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos referenciados en la lista adjunta (2 páginas), 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.*

*Estos dispositivos se clasifican en la categoría "otro dispositivo", ya que no pertenecen a la lista A ni a la lista B del anexo II, tampoco a la clase de autodiagnóstico.*

*Esta declaración se basa en el contenido de cada expediente técnico y está respaldado por la certificación de nuestro sistema de calidad según la norma NF EN ISO 13485 : 2016 (Certificación válida hasta el 27 de Julio 2023).*

Sées, le 29 juillet 2020

**Valérie LAMBERT,**

Responsable des Affaires Réglementaires

*Regulatory Affairs Manager*

*Responsable de los Asuntos Reglamentarios*

**Cécile GOUBAULT,**

Directeur Général Délégué

*Managing Director*

*Directora General*

REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCES - REFERENCIAS	Code GMDN
<b>Metabolites divers / Miscellaneous metabolites</b>		
ALBUMIN	ALBU-0600/0700/0250	53597
ALBUMIN ENVOY	ALBU-0850	
BILIRUBIN DIRECT 4+1	BIDI-0600/0250	53233
BILIRUBIN TOTAL 4+1	BITO-0600/0250	53229
BILIRUBIN TOTAL & DIRECT 4+1	BITD-0600	53229/53233
CREATININE ENVOY	CRSL-0850	53250
CREATININE JAFFE	CRCO-0600/0700	53251
CREATININE PAP SL	CRSL-0630/0250	53250
DIRECT BILIRUBIN ENVOY	BIDV-0850	53233
GLUCOSE ENVOY	GPSL-0850	
GLUCOSE HK SL	GHSL-0600/0250	53301
GLUCOSE PAP SL	GPSL-0507/0500/0707/0700/0250/0455/0497	
LACTATE	LACT-0100	53342
MICROPROTEIN PLUS	PRTU-0600/0250	53481
PHOSPHORUS	PHOS-0600/0230	
PHOSPHORUS ENVOY	PHOS-0850	59123
TOTAL BILIRUBIN ENVOY	BITV-0850	53229
TOTAL PROTEIN ENVOY	PROB-0850	
TOTAL PROTEIN PLUS	PROB-0600/0700/0250	53985
UREA ENVOY	URSL-0850	
UREA UV SL	URSL-0407/04270420/0500/0507/0250/0455	53587
URIC ACID ENVOY	AUVD-0850	
URIC ACID MONO SL	AUML-0497/0427/0420/0500/0507/0707/0250	
URIC ACID SL	AUSL-0250	53583
<b>Enzymes / Enzymes</b>		
ALP (DEA) SL	PASL-0400/0420/0230	
ALP ENVOY	PIVD-0850	52928
ALP IFCC	ALPI-0230	
ALT ENVOY	ALSL-0850	
ALT/GPT 4+1 SL	ALSL-0410/0430/0510/0250/0455	52923
AMYLASE ENVOY	AMSL-0850	
AMYLASE SL	AMSL-0390/0400/0230	52940
AST ENVOY	ASVD-0850	
AST/GOT 4+1 SL	ASSL-0410/0430/0510/0250/0455	52954
CHOLINESTERASE	CHES-0053	52971
CK ENVOY	CKSL-0850	53003
CK-MB ENVOY	CMSL-0850	
CK-MB SL	CMSL-0410/0430/0230	52994
CK NAC SL	CKSL-0410/0430/0230	53003
GAMMA-GT PLUS SL	GISL-0400/0420/0250	
GGT ENVOY	GISL-0850	53027
LDH ENVOY	LLSL-0850	
LDH-L SL	LLSL-0400/0420/0230	53072
LIPASE ENVOY	LPSL-0850	
LIPASE SL	LPSL-0230	53108
<b>Electrolytes - Oligo-éléments / Electrolytes - Trace-elements</b>		
CALCIUM ARSENATO	CALA-0600/0250	
CALCIUM ENVOY	CALA-0850	45789
CHLORIDE	CHLO-0600/0250	60037
IRON ENVOY	FEFE-0850	
IRON FERENE	FEFE-0230/0600	54758
MAGNESIUM ENVOY	MAGX-0850	
MAGNESIUM XB	MGXB-0250/0600	
MAGNESIUM XYLIDYL	MAGX-0230/0600	46795
<b>Lipides / Lipids</b>		
CHOLESTEROL ENVOY	CHSL-0850	
CHOLESTEROL SL	CHSL-0507/0500/0700/0707/0250/0455/0497	53359
CHOLESTEROL HDL SL 2G	HDLL-0230/0380/0390	53391
CHOLESTEROL LDL SL 2G	LDLL-0230/0380/0390	53395
HDL CHOLESTEROL	CHDL-0250/0600	
HDL CHOLESTEROL ENVOY	HDLL-0850	53391
LDL CHOLESTEROL	CLDL-0250	
LDL CHOLESTEROL ENVOY	LDLL-0850	53395
TRIGLYCERIDES ENVOY	TGML-0850	
TRIGLYCERIDES MONO SL NEW	TGML-0427/0425/0515/0700/0517/0707/0497	53460
TRIGLYCERIDES SL	TGML-0250/0455	
<b>Contrôles-Calibrants-Standards / Controls-Calibrators-Standards</b>		
CHOLESTEROL HDL 2G CALIBRATOR	HDLL-0011/0041	44696
CHOLESTEROL LDL 2G CALIBRATOR	LDLL-0011/0041	41728
CHOLESTEROL Standard 200 mg/dL	CHOL-0055	44698
CK-MB CONTROL	CKMB-0900	44693
ELICAL 2	CALI-0550	47868
ELITROL I	CONT-0060	
ELITROL II	CONT-0160	47869
GLUCOSE Standard 100 mg/dL	GLUP-0055	41818
HDL LDL CALIBRATOR	HLCA-0041	47868
ISE CONTROL I	ISCT-0046	
ISE CONTROL II	ISCT-0047	47869
MICROPROTEIN PLUS Standard 100 mg/dL	PRTU-0022	53482
TRIGLYCERIDES Standard 200 mg/dL	TRIG-0055	44702
UREA Standard 50 mg/dL	URUV-0055	53588
URIC ACID Standard 6 mg/dL	ACUR-0055	44704

REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCES - REFERENCIAS	Code GMDN
<b>Protéines spécifiques / Specific proteins</b>		
ANTI-STREPTOLYSIN O	ASLO-0250	59055
CRP IP	ICRP-0400	53705
CRP IP CALIBRATOR SET	ICRP-0043	41838
CRP IP CONTROL I	ICRP-0046	41839
CRP IP CONTROL II	ICRP-0047	
CRP WR	CRPW-0230	53705
CRP WR CALIBRATOR SET	CRPW-0043	41838
CRP WR CONTROL	CRPW-0045	41839
CRP WR ENVOY	CRPW-0850	53705
FERRITIN	IFRT-0230	53718
FERRITIN CALIBRATOR	IFRT-0042	41927
HAPTOGLOBIN IP	IHAP-0400	53737
HbA1c	HBAC-0240	59090
HbA1c CALIBRATOR SET	HBAC-0043	53315
HbA1c CONTROL L + H	HBAC-0049	44435
IgA IP	IIGA-0400	53760
IgG IP	IIGG-0400	53787
IgM IP	IIGM-0400	53795
μALBUMIN IP	IMAL-0400	53475
μALBUMIN IP CALIBRATOR SET	IMAL-0043	53477
μALBUMIN IP CONTROL I	IMAL-0046	
μALBUMIN IP CONTROL II	IMAL-0047	53478
OROSOMUCOID IP	IORT-0400	53606
PREALBUMIN IP	IPAL-0400	53957
PROTEIN IP CALIBRATOR SET	IPRO-0043	53593
RF CALIBRATOR	IRFA-0042	42230
RHEUMATOID FACTOR	IRFA-0230	55111
RHEUMATOLOGY CONTROL I	IRCT-0046	47869
RHEUMATOLOGY CONTROL II	IRCT-0047	
TRANSFERRIN IP	ITRF-0400	59041
<b>Vitamines/Vitamins</b>		
VITAMIN D	VITD-0250	54476
VITAMIN D CALIBRATOR SET	VITD-0043	54474
VITAMIN D CONTROL SET	VITD-0049	54475
<b>ISE Solutions pour électrodes selectives d'ions / ISE Solutions for ion-selective electrodes</b>		
ISE BASELINE SOLUTION ENVOY	ISBA-0850	59238
ISE CALIBRATORS	ISCA-0250	52867
ISE CALIBRATOR ENVOY	ISCV-0850	
ISE CLEANER/CONDITIONER	ISCC-0280	59058
ISE DILUENT	ISDI-0250	58237
ISE DILUENT ENVOY	ISDV-0850	
ISE REFERENCE SOLUTION	ISRS-0800	59238
ISE REFERENCE SOLUTION ENVOY	ISRS-0850	
<b>Solutions de lavage pour les équipements ELITech Clinical Systems / Cleaning solutions for ELITech Clinical Systems Equipments</b>		
ACID SOLUTION for ELITech Clinical Systems Analyzers	SLHC-5900	59058
SYSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers	SLNA-5900	59058
SYSTEM SOLUTION	SLSY-5905	58236
SYSTEM SOLUTION for ELITech Clinical Systems Analyzers	SLSY-5900	
<b>Tests d'agglutination / Agglutination tests</b>		
CRP LATEX	LXCR-0112	53707



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Register

# Certificate of Approval

This is to certify that the Management System of:

**ELITechGroup B.V.**

**Van Rensselaerweg 4, 6956 AV Spankeren, Netherlands**

has been approved by LRQA to the following standards:

**ISO 13485:2016**

David Derrick - Area Operations Manager UK & Ireland

Issued By: Lloyd's Register Quality Assurance Limited

This certificate is valid only in association with the certificate schedule bearing the same number on which the locations applicable to this approval are listed.

**Current Issue Date:** 22 June 2018

**Original Approvals:**

**Expiry Date:** 21 June 2021

**ISO 13485 21 November 2016**

**Certificate Identity Number:** 10093739

**Certificate Approval Number:** LRQ 00000428

**Product Approval Number:** ISO 13485 – 0016037

The scope of this approval is applicable to:

**Design, development, manufacture and distribution of clinical chemistry analyzers and erythrocyte sedimentation rate analyzers and tubes for the in vitro diagnostic investigation of samples of human origin..**



001



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# Certificate Schedule

Certificate Identity Number: 10093739

## Location

## Activities

Van Rensselaerweg 4, 6956 AV Spankeren,  
Netherlands

ISO 13485:2016

Design, development, manufacture and distribution of clinical chemistry analyzers and erythrocyte sedimentation rate analyzers and tubes for the in vitro diagnostic investigation of samples of human origin.

Kanaaldijk 90, 6956 AX Spankeren, Netherlands

ISO 13485:2016

Warehousing of parts, finished instruments and erythrocyte sedimentation rate tubes.



001



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	HBe Ag&Ab CODE: HBE.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST A
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY	AEMPS – n° 0318
(EC) CERTIFICATE(S)	<ul style="list-style-type: none"><li>• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318</li><li>• DESIGN CERTIFICATE N° 2004 03 0425 ED RELEASED BY EC NOTIFIED BODY N° 0318</li><li>• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318</li></ul>

PLACE & DATE OF FIRST ISSUE	MILANO – APRIL 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 12/2013

# Declaration of Conformity



HL-7-0137DC DOI 2015/07 (7)

**In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.**

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5186	Routine Control N	30590

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson

Title: Managing Director

Signed:

Date: 28 Jul 2015

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United Kingdom

# Declaration of Conformity

**helena**  
Biosciences Europe

HL-7-0138DC DOI 2015/07 (7)

**In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.**

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5187	Routine Control A	30590

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson

Title: Managing Director

Signed:



Date: 28 Jul 2015

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United Kingdom

# Declaration of Conformity



HL-7-0163DC DOI 2015/08 (9)

**In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.**

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5265	Thromboplastin LI	55983
5265H	Thromboplastin LI	55983

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson

Title: Managing Director

Signed:

Date: 06 Aug 2015

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United Kingdom

# Declaration of Conformity



HL-7-0286DC DOI 2015/08 (5)

**In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.**

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5558	APTT Si L Minus	55981

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson

Title: Managing Director

Signed:

Date: 11 Aug 2015

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# Declaration of Conformity



HL-7-0567DC DOI 2015/08 (1)

**In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.**

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5376R	Clauss Fibrinogen 100	55997

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson

Title: Managing Director

Signed:

Date: 12 Aug 2015

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United Kingdom



Dia.Pro  
Diagnostic  
BioProbes

# EC DECLARATION OF CONFORMITY

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

<b>ISO CERTIFICATE</b>	UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY AEMPS (AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS)
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<b>PLACE &amp; DATE OF FIRST ISSUE</b>	MILANO – MARCH 2004
<b>PLACE &amp; DATE OF CURRENT ISSUE</b>	SESTO SAN GIOVANNI (MI) – MARCH 2019
<b>SIGNATURE</b> <b>Legal Representative</b> <b>Dr.ssa Fiorenza Scozzesi</b>	

Rev: 05/2018