

bsi.



By Royal Charter

Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016 & EN ISO 13485:2016

This is to certify that:

Helena Laboratories (UK) Ltd
trading as Helena Biosciences Europe
Queensway South
Team Valley Trading Estate
Gateshead
Tyne and Wear
NE11 0SD
United Kingdom

Holds Certificate Number:

MD 69326

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 & EN ISO 13485:2016 for the following scope:

The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.

For and on behalf of BSI:

Stewart Brain

Stewart Brain, Head of Compliance & Risk - Medical Devices

Original Registration Date: 2002-10-25

Latest Revision Date: 2018-11-28

Effective Date: 2018-04-14

Expiry Date: 2021-04-13

Page: 1 of 2



...making excellence a habit™

Certificate No: **MD 69326**

Location

Helena Laboratories (UK) Ltd
trading as Helena Biosciences Europe
Sunderland Enterprise Park
Colina Avenue
Sunderland
SR5 3XB
United Kingdom

Registered Activities

The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.

Location

Helena Laboratories (UK) Ltd
trading as Helena Biosciences Europe
Queensway South
Team Valley Trading Estate
Gateshead
Tyne and Wear
NE11 0SD
United Kingdom

The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.

Original Registration Date: 2002-10-25
Latest Revision Date: 2018-11-28

Effective Date: 2018-04-14
Expiry Date: 2021-04-13

Page: 2 of 2



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BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W6 4AL, UK.
A Member of the BSI Group of Companies.

CERTIFICAT OF REGISTRATION N° 10462 rev. 6

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

ELITTECH CLINICAL SYSTEMS SAS
Zone Industrielle
61500 SEES FRANCE

pour les activités
for the activities

Conception, production, contrôle et commercialisation de réactifs de chimie clinique
destinés aux laboratoires d'analyses de biologie.
Validation de la combinaison réactifs et automates.

Design, production, control and sales of clinical chemistry reagents
intended to be used by clinical laboratories.
Validation of the combination reagents and instruments.

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

ELITTECH 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

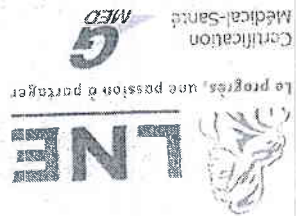
Début de validité / Effective date : July 28th, 2017 (included)
Valable jusqu'au / Expiry date : July 27th, 2020 (included)
Etabli le / Issued on : July 27th, 2017

On behalf of the Certification Director
Cécile VAUGELADE
G-MED Certification Division Manager



LNE N° 10462-6
Ce certificat est délivré selon les règles de certification G-MED / This certificate is issued according to the rules of G-MED certification
Renouvellement le certificat 10462-5

Laboratoire national de métrologie et d'essais • Etablissement public à caractère industriel et commercial
LNE/G-MED • Organisme notifié par l'autorité compétente française en matière de dispositifs médicaux n° 0459
1, rue Gaston Boissier - 75724 Paris Cedex 15 • Tél. : 01 40 43 37 00 • Fax : 01 40 43 37 37 • www.lne.fr • www.gmed.fr



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DECLARATION DE CONFORMITE CE

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

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

(Voir liste ci-jointe).

DECLARATION OF EC CONFORMITY

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

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

(See attached list).

DECLARACIÓN CE DE CONFORMIDAD

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

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

(Ver lista adjunta).

Sees, le 18 Mai 2018

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

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Cécile GOUBAULT,
Directeur Général Délégué
Managing Director
Directora General



GRUPE 1 - METABOLITES DIVERS GROUP 1 - MISCELLANEOUS METABOLITES GRUPO 1 - METABOLITOS VARIOS

DESIGNATION DU REACTIF/ REAGENT DESIGNATION/ DESIGNACIÓN DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO	Code GMDN/ GMDN Code/ Codigo GMDN
ALBUMIN	ALBU-0600/0700/0250	DOS-CE-ALBU_RASid	53597
ALBUMIN ENVOY	BIDJ-0600/0250		53233
BILIRUBIN DIRECT 4+1	BITO-0600/0250	DOS-CE-BILU 4+1	53229
BILIRUBIN TOTAL 4+1	BITO-0600/0250		53229/53233
CREATININE ENVOY	CRSL-0850	DOS-CE-CRSL	53250
CREATININE JACFE	CRCO-0600/0700	DOS-CE-CRCO_RASid	53251
CREATININE PAP SL	CRSL-0630/0250	DOS-CE-CRSL	53250
DIRECT BILIRUBIN ENVOY	BIDV-0850	DOS-CE-BIDV	53233
GLUCOSE ENVOY	GPSL-0850	DOS-CE-GPSSL_RASid	
GLUCOSE IR SL	GHSI-0600/0250	DOS-CE-GHSI	53301
GLUCOSE PAP SL	GPSSL-0495/0500/0700/ 0507/0707/0250/0455/0497	DOS-CE-GPSSL_RASid	
HEMOGLOBIN	HBAO-0400	DOS-CE-HEMO	32430
IRON TIBC	FECA-0050	DOS-CE-TIBC	53904
LACTATE	LACT-0100	DOS-CE-LACT	53442
MICROPROTEIN PLUS	PRU-0600/0250	DOS-CE-PRU_RASid	53481
PHOSPHORUS	PHOS-0600/0230	DOS-CE-PRU_RASid	59123
PHOSPHORUS ENVOY	PHOS-0850	DOS-CE-PRU_RASid	59123
TOTAL BILIRUBIN ENVOY	BITV-0850	DOS-CE-BITV	53229
TOTAL PROTEIN	PRTB-0600	DOS-CE-PRTB	
TOTAL PROTEIN ENVOY	PROB-0850	DOS-CE-PROB_RASid	53985
TOTAL PROTEIN PLUS	PROB-0600/0700/0250		
UREA ENVOY	URSL-0850	DOS-CE-URSL_RASid	
UREA 64	URU-0400	DOS-CE-URU	53587
UREA UT SL	URSL-0400/0420/0500 0407/0427/0507/0250/0455	DOS-CE-URSL_RASid	
URIC ACID	ACUR-0200/0400	DOS-CE-ACUR	
URIC ACID ENVOY	AUVD-0850	DOS-CE-AUVD	53583
URIC ACID MQUO SL	AUML-0420/0500/0700/ 0427/0497/0507/0707/0250	DOS-CE-AUML_RASid	
URIC ACID SL	AUML-0250	DOS-CE-AUML	

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DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone Industrielle 61500 Sées France, déclarons sous notre seule responsabilité que les réactifs appartenant au groupe 2 « ENZYMES », référencés dans la liste ci-jointe, sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.

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

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 Sées France, hereby certify, under our own responsibility, that the reagents belonging to Group 2, "ENZYMES", such as listed hereon, conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to *in vitro* diagnostic medical devices and to the public health code.

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

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 Sées France, declaramos bajo nuestra única responsabilidad que los reactivos pertenecientes al grupo 2 : "ENZIMAS", referenciados en la lista adjunta, son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico *in vitro* y el código de salud pública.

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

Sées, le 28 Juillet 2017

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

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Cécile COUBAULT,
Directeur Général Délégué
Managing Director
Directora General



GRUPE 2 - ENZYMES GROUP 2 - ENZYMES GRUPO 2 - ENZIMAS

DESIGNATION DU REACTIF / REACTIF DESIGNATION/ DESIGNACIÓN DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO CE	Code GMDN/ GMDN Code/ Código GMDN
ALP IDEAL SL	PASL-0400/0420/0230	DOS-CE-PASL	52928
ALP ENVOY	PIVD-0850	DOS-CE-PIVD	
ALT ENVOY	ALSL-0850	DOS-CE-ALSL 4+1	52923
ALT /GPT	ALAT-0200/0400	DOS-CE-ALAT	
ALT /GPT +1 SL	ALSL-0410/0430/0510/0250/0455	DOS-CE-ALSL 4+1	
AMYLASE ENVOY	AMSL-0850	DOS-CE-AMSL	52940
AMYLASE SL	AMSL-0390/0400/0230	DOS-CE-AMSL	
AST ENVOY	ASVD-0850	DOS-CE-ASVD	
AST /GOT	ASAT-0200/0400	DOS-CE-ASAT	52954
AST /GOT +1 SL	ASSL-0410/0430/0510/0250/0455	DOS-CE-ASSL 4+1	
CHOLINESTERASE	CHES-0053	DOS-CE-CHES	52971
CK ENVOY	CKSL-0850	DOS-CE-CKSL	53003
CK-MB	CKMB-0030	DOS-CE-CKMB	
CK-MB ENVOY	CKMSL-0850	DOS-CE-CKMSL	52994
CK-MB SL	CKMSL-0410/0430/0230	DOS-CE-CKMSL	
CK NAC	CKNA-0030	DOS-CE-CKNA	53003
CK NAC SL	CKSL-0410/0430/0230	DOS-CE-CKSL	
CK-MMA GT SL PLUS	GISL-0400/0420/0500/0250	DOS-CE-CKSL	
CK-MMA ENVOY	GISL-0850	DOS-CE-CKSL	53027
LDH ENVOY	LLSL-0850	DOS-CE-LLSL	
LDH SL	LLSL-0400/0420/0230	DOS-CE-LLSL	53072
LDH-P	LDHP-0030	DOS-CE-LDHP	
LDH-P ENVOY	LDPSL-0850	DOS-CE-LDHP	
LDH-P SL	LDPSL-0230	DOS-CE-LDPSL	53108

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DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone Industrielle 61500 SEES France, déclarons sous notre seule responsabilité que les réactifs appartenant au groupe 3 «ELECTROLYTES/OLIGO-ELEMENTS», référencés dans la liste ci-jointe, sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.
Cette déclaration est basée sur le contenu de chaque dossier technique DOS-CE-XXXX et s'appuie sur la certification de notre système qualité selon la norme NF EN ISO 13485 : 2016 (Certification valable jusqu'au 27 juillet 2020).
(Voir liste ci-jointe).

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents belonging to Group 3 "ELECTROLYTES/TRACE-ELEMENTS", such as listed hereto, conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to *in vitro* diagnostic medical devices and to the public health code.
This declaration is based on the contents of each DOS-CE-XXXX technical file and is supported by the certification of our quality system according to the standard NF EN ISO 13485 : 2016 (Certification valid until July 27th, 2020).
(See attached list).

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos pertenecientes al grupo 3 "ELECTROLITOS/OLIGO-ELEMENTOS", referenciados en la lista adjunta, son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico *in vitro* y el código de salud pública.
Esta declaración se basa en el contenido de cada DOS-CE-XXXX técnica y está respaldado por la certificación de nuestro sistema de calidad según la norma NF EN ISO 13485 : 2016 (Certificación válida hasta el 27 de Julio 2020).
(Ver lista adjunta)

Sees, le 29 juin 2018

Valérie GOURDON,

Responsable des Affaires Réglementaires
Regulatory Affairs Manager
Responsable de los Asuntos Reglamentarios

Valérie Gourdon

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GRUPE 3 – ELECTROLYTES / OLIGO-ELEMENTS GROUP 3 – ELECTROLYTES / TRACE-ELEMENTS GRUPO 3 – ELECTROLITOS / OLIGO-ELEMENTOS

DESIGNATION DU REACTIF/ REAGENT DESIGNATION/ DESIGNACIÓN DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO CE	Code GMDN/ GMDN Code/ Código GMDN
CALCIUM ARSENAZO	CALA-0600/0250	DOS-CE-CALA_R&Sid	45789
CALCIUM ENVOY	CALA-0850		
CHLORIDE	CHLO-0600/0250	DOS-CE-CHLO_R&Sid	60037
IRON CHROMAZUROL	FECA-0600	DOS-CE-FECA	
IRON ENVOY	FEFE-0850	DOS-CE-FEFE_R&Sid	54758
IRON FERRENE	FEFE-0230/0600		
IRON FERROZINE	FEFR-0600	DOS-CE-FEFR_R&Sid	
MAGNESIUM CALMAGITE	MAGN-0600	DOS-CE-MAGN_R&Sid	
MAGNESIUM ENVOY	MAGX-0850		
MAGNESIUM XYLIDYL	MAGX-0230/0600	DOS-CE-MAGX-R&Sid	46795

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DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone Industrielle 61500 SEES France, déclarons sous notre seule responsabilité que les réactifs appartenant au groupe 4 «LIPIDES », référencés dans la liste ci-jointe, sont conformes aux exigences essentielles des annexes I et II de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.

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

(Voir liste ci-jointe).

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents belonging to Group 4 "LIPIDS", such as listed here, conform to the essential requirements of appendices I and II of European Directive 98/79/EC, relating to *in vitro* diagnostic medical devices and to the public health code.

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

(See attached list).

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos pertenecientes al grupo 4 "LIPIDOS ", referenciados en la lista adjunta, son conformes con los requisitos esenciales de los anexos I y II de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico *in vitro* y el código de salud pública.

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

(Ver lista adjunta)

Sees, le 28 Juillet 2017

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



GROUPE 4 – LIPIDES GROUP 4 – LIPIDS GRUPO 4 – LÍPIDOS

DESIGNATION DU REACTIF/ REAGENT DESIGNATION/ DESIGNACION DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO CE	Code GMDN/ GMDN Code/ Codigo GMDN
CHOLESTEROL	CHOL-0220/0420	DOS-CE-CHOL	
CHOLESTEROL ENVOY	CHSL-0850		53359
CHOLESTEROL SL	CHSL-0495/0500/0700/ 0507/0707/0250/0455/0497	DOS-CE-CHSL	
CHOLESTEROL HDL SL 2G	HDLL-0230/0380/0390	DOS-CE-HDLL	53391
CHOLESTEROL LDL SL 2G	LDLL-0230/0380/0390	DOS-CE-LDLL	53395
HDL CHOLESTEROL	HDLC-0060	DOS-CE-HDLC	53391
HDL CHOLESTEROL ENVOY	HDLL-0850	DOS-CE-HDLL	
LDL CHOLESTEROL ENVOY	LDLL-0850	DOS-CE-LDLL	53395
TRIGLYCERIDES	TRIG-0200/0400	DOS-CE-TRIG	
TRIGLYCERIDES ENVOY	TGML-0850		53460
TRIGLYCERIDES MONO SL NEW	TGML-0425/0495/0515/ 0700/0427/0517/0707/0497	DOS-CE-TGMLN	
TRIGLYCERIDES SL	TGML-0250/0455		

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DECLARATION DE CONFORMITE CE

Nous, ELITech Clinical Systems SAS, zone industrielle 61500 SÉES France, déclarons sous notre seule responsabilité que les réactifs appartenant au groupe 5 «CONTROLES/ CALIBRANTS/ STANDARDS», référencés dans la liste ci-jointe, sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.

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

(Voir liste ci-jointe).

DECLARATION OF EC CONFORMITY

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents belonging to group 5, "CONTROLS/ CALIBRATORS/ STANDARDS", such as listed hereto, conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to *in vitro* diagnostic medical devices and to the public health code.

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

(See attached list).

DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos pertenecientes al grupo 5 "CONTROLES/ CALIBRADORES/ ESTÁNDARES", referenciados en la lista adjunta, son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico *in vitro* y el código de salud pública.

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

(Ver lista adjunta)

Sées, le 28 Juillet 2017

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

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Cécile GOURBAUT,
Directeur Général Délégué
Managing Director
Directora General

GRUPE 5 – CONTROLES/CALIBRANTS/STANDARDS GROUP 5 – CONTROLS/CALIBRATORS/STANDARDS GRUPO 5 – CONTROLES/CALIBRADORES/ESTÁNDARES

DESIGNATION DU REACTIF/ REAGENT DESIGNATION/ DESIGNACION DE REACTIVO	REFERENCES/ REFERENCIAS	NOM DU DOSSIER CE/ EC FILE NAME/ NOMBRE DEL ARCHIVO CE	Code GMDN/ GMDN Code/ Código GMDN
CHOLESTEROL HDL 2G CALIBRATOR	HDL-0011/0041	DOS-CE-HDL-CAL	44696
CHOLESTEROL LDL 2G CALIBRATOR	LDL-0011/0041	DOS-CE-LDL-CAL	41728
CHOLESTEROL Standard 200 mg/dL	CHOL-0055	DOS-CE-CHOL200	44698
CK-MB CONTROL	CKMB-0900	DOS-CE-CKMB-CT	44693
CREATININE Standard 2 mg/dL	CREN-0055	DOS-CE-CREN2	44700
ELICAL 2	CAL-0550	DOS-CE-CAL12	47868
ELITROL I	CONT-0060	DOS-CE-ELT I	47869
ELITROL II	CONT-0160	DOS-CE-ELT II	47869
GLUCOSE Standard 100 mg/dL	GLUP-0055	DOS-CE-GLUP100	41818
ISE CONTROL I	ISCT-0046	DOS-CE-ISCT	47869
ISE CONTROL II	ISCT-0047	DOS-CE-ISCT	47869
MICROPROTEIN PLUS	PRTU-0022	DOS-CE-PRTU100	53482
Standard 100 mg/dL			
TRIGLYCERIDES Standard 200 mg/dL	TRIG-0055	DOS-CE-TRIG200	44702
UREA Standard 50 mg/dL	URUV-0055	DOS-CE-URUV50	53588
URIC ACID Standard 6 mg/dL	ACUR-0055	DOS-CE-ACUR6	44704

ELITech Clinical Systems SAS

Zone Industrielle
61500 SÉES - France
Tél : +33 (0)2 33 81 21 00 Fax : +33 (0)2 22 28 77 51
www.elitechgroup.com

СИСТЕМА СЕРТИФИКАЦИИ
ФЕДЕРАЛЬНОЕ АГЕНТСТВО ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И
МЕТРОЛОГИИ

СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ

«СМК СТАНДАРТ»

Per. № РОСС RU.31060.04ЖКЖЮ0

Орган по сертификации:

РФТ № SMK STANDARD.TU.0005

Общество с ограниченной ответственностью

«МЕЖДУНАРОДНЫЙ ЦЕНТР СЕРТИФИКАЦИИ»

Адрес: 190020, г. Санкт-Петербург, наб. Обводного канала, д. 138, корпус 1, офис 421

тел +7 (812) 438-76-71 standard@iso-smk.ru

полнота сертификата проверяйте в реестре на сайте <http://www.iso-smk.ru>

СЕРТИФИКАТ СООТВЕТСТВИЯ

№ ST.RU.0001.M0013380

выдан

Обществу с ограниченной ответственностью «Атат-Мед»

Адрес: 105173, г. Москва, ул. Главная, д.6, кв.12

ИНН 7719187311 ОГРН 1037739078970

Дата выдачи: 26.01.2018 г. Срок действия до: 26.01.2021 г.

Настоящий сертификат удостоверяет:

Идентификационные Системы менеджмента качества

Системы менеджмента качества применительно к работам

составляющим часть сертификации

(приложение является неотъемлемой частью сертификата)

СООТВЕТСТВУЕТ ТРЕБОВАНИЮ ГОСТ ISO 13485-2011 (EN ISO 13485:2003)

Руководитель органа

Концев В. В.

Эксперт

Лыкина О. В.





Тынарепа О. В.

Зксепт



Копие В. В.

Вукановиче, Опрана

Разработка, производство и продажа медицинских изделий для in vitro диагностики: реанитов и наборов реанитов для клинической диагностики, а также реанитов и реанитов.

Область сертификации системы менеджмента качества:

к сертификату соответствия № ST.RU.0001.M0013380

ПРИЛОЖЕНИЕ №1

МЕТОДОЛГИИ

СИСТЕМА СЕРТИФИКАЦИИ
ФЕДЕРАЛЬНОЕ АГЕНТСТВО ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И



ООО "АГАТ-МЕД"
105173, Москва, ул. Главная 6-12
многоканальный т/ф. 777-41-92
факс/авт. (495) 741-25-19;
т/ф. (499) 780-97-48/84.
E-mail: agat@agat.ru, http://www.agat.ru

ПАСПОРТ

Набор реагентов для определения гемоглобина в крови
темнотиминным методом

«Темнотимин-АГАТ»

(600 опр. x 5 мл)

Серия 17/960918 Дата выпуска 09.2018 Годен до 50000 10.2020

Наименование показателя	Характеристика и норма по ТУ 9398-280-11498242-00	Результаты анализа
-------------------------	--	--------------------

1. Внешний вид	1.1. Трансформирующий реагент 1.2. Ацетонцианидурин 1.3. Катрибировочный раствор гемоглобина	Лорошок желто-оранжевого цвета или смесь белых и оранжевых кристаллов бесцветная прозрачная жидкость прозрачная жидкость темно-красного цвета
----------------	--	---

2. Технические характеристики	2.1 Тест на соответствие катрибрату контрольному лабораторному гемоглобину 120 г/л, отклонение в %, не более 2.2 Чувствительность, г/л, не более 3. Показатели правильности определения	2 10	Соответствует	Соответствует
3.1 Тест на "линейность" в диапазоне концентраций от 20 до 200 г/л, отклонение в %, не более 3.2 Тест на "открытие", отклонение в %, не более 3.3 Коэффициент вариации результатов определения, %, не более 3.4 Допустимый разброс результатов при параллельных определениях одной пробы разными катрибратом одной серии, %, не более 3.5 Время выхода оптической плотности на устойчивые показатели после добавления анализируемой пробы, мин, не более	20 2 2 2 2	Соответствует	Соответствует	Соответствует

Заключение ОКК ООО «Агат-Мед»:
Серия 17/960918 соответствует требованиям ТУ 9398-280-11498242-00

Начальник ОКК ООО «АГАТ-МЕД» Галуза В.В.
«1» сентября 2018 г.



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРПЛОТООХРАНЕНИЯ
(РОСЗАРПЛАТНАДЗОР)



РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДАНИЕ

от 25 ноября 2011 года № ФСР 2011/12395

На медицинское издание
Набор рефертов для определения гемоглобина в крови темнотемпературостойким
методом «Смолтобин-Атат» по ТУ 9398-280-11498242-00

Настоящее регистрационное удостоверение выдано
Обществу с ограниченной ответственностью "Атат-Мед" (ООО "Атат-Мед"),
Россия, 105173, Москва, поселок Восточный, ул. Гавриная, д. 6, кв. 12

Производитель
Общество с ограниченной ответственностью "Атат-Мед" (ООО "Атат-Мед"),
Россия, 105173, Москва, поселок Восточный, ул. Гавриная, д. 6, кв. 12

Место производства медицинского изделия
105173, Москва, поселок Восточный, ул. Гавриная, д. 6, кв. 12
Идентификационный номер № 41794 от 19.10.2011

Вид медицинского изделия -

Класс потенциального риска применения медицинского изделия I
Код Общероссийского классификатора продукции для медицинского изделия 93 9816
Настоящее регистрационное удостоверение имеет приложение на 1 листе
Идентификационный номер от 25 ноября 2011 года № 7750-17/11



и приказом от 10 декабря 2013 года № 7123-17/13 о замене
получено к обработке на территории Российской Федерации.

Взно руководителем Федеральной службы
по патентам в сфере здравоохранения

0005897

ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВОСХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

ПРИЛОЖЕНИЕ

К РЕГИСТРАЦИОННОМУ УДОСТОВЕРЕНИЮ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 25 ноября 2011 года № ФСР 2011/12395

Лист 1

На медицинское изделие

Набор реагентов для определения гемоглобина в крови гемодиализными
методом «Гемоглобин-Атат» по ТУ 9398-280-11498242-00
1. Трансформирующий реагент (сухая смесь 200 мЛ) - 3 упаковки;
2. Аутофлуоресцентный реагент (по 0,5 мЛ);
3. Калибровочный раствор гемоглобина с концентрацией 120 г/Л - 1 флакон (2 мЛ).



З

Приказом от 10 декабря 2013 года № 7123-1п/13 о замене/дополнении и
регистрации Российской Федерации.

Вручено руководителю Департамента
по надзору в сфере здравоохранения

М.А. Мухомов

0005004

CERTIFICATE OF REGISTRATION



Lorne Laboratories Ltd

Unit 1 Cutbush Park Industrial Estate

Danehill

Lower Earley

Berkshire RG6 4UT UNITED KINGDOM

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

ISO 13485:2016

EN ISO 13485:2016

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

Authorized by

Michael J. Windler

Michael J. Windler, P.E.

Manager of Global Regulatory Service

Distinguished Member of the Technical Staff

UL Life and Health Sciences

UL LLC

Check Certificate
Status: here



File Number A12241
Certificate 1458.180626
Initial Issue June 26, 2018

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

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



00-MB-50043 Issue 15.0

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





EC DECLARATION OF CONFORMITY

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

Product name	CRP Latex kit
Catalogue number	850100A

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

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

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

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

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

Eddy Velthuis
Technical Director



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

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





EC DECLARATION OF CONFORMITY

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

Product Name	Catalogue Number
LISS Ready For Use	470020 470250 470025

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

This product has been manufactured by Lorne Laboratories Ltd at their facilities which are located at:

Address line 1: Unit 1 Cutbush Park Industrial Estate
Address line 2: Danehill
City: Lower Earley
County: Berkshire
Postal code: RG6 4UT
Country: United Kingdom
Telephone number: +44-(0)118 921 2264
Fax number: +44-(0)118 986 4518
Website: www.lornelabs.com
DUNS Number: 732670703

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

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

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



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



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



This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 10 April 2019.

Eddy Velthuis
Technical Director



ISO 13485:2015
GD 19400001, 02/01/2020

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

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

CERTIFICATE OF REGISTRATION

This is to certify that the quality management system of:

Medica Corporation

Main Site: 5 Oak Park Drive

Bedford, Massachusetts 01730 United States

has been assessed by Intertek as conforming to the requirements of:

ISO 13485:2016

The quality management system is applicable to:

The Design, Development, Manufacture, Service, Distribution of in-vitro diagnostic medical devices, in-vitro diagnostic test kits, in-vitro diagnostic reagents, in-vitro diagnostic analyzers/software used in the diagnosis and management of cancer, immune status, disease status, autoimmune status, cardiac markers, protein metabolism, endocrine disorders, blood analytes, urinalysis, blood gases.

Certificate Number: 0082581-01

Initial Certification Date: 2009-04-17

Certificate Issue Date: 2019-01-01

Certificate Expiry Date: 2021-04-16



Calin Ividoveanu
President
Intertek Testing Services NA Ltd.
1829, 32nd Avenue, Lachine, QC H8T 3J1,
Canada



MEDICA

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

Products For Health Care

Declaration of Conformity CE

Product Name:

EasyLyte and accessories per attachment

EasyElectrolyte and accessories per attachment

EasyStat and accessories per attachment

EasyBloodGas and accessories per attachment

Manufacturer

Medica Corporation
5 Oak Park Drive, Bedford, Massachusetts, 01730, USA

Representative

EC REP Emergo Europe, Molensstraat 15
NL-2513 BH 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 in conformity with the Annex III, essential requirements and provisions of Council Directive: 98/79/EC

Place and Date: Bedford, Massachusetts, USA, March 1, 2012

Signature:

Photios Makris

Name: Photios Makris

Title: Director of Regulatory Affairs



Model/Type:

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

EasyElectrolyte Na/K/Cl, Na/K/Li

pH/pCO2/pO2/Na/K/Ca/Hct, pH/pCO2/pO2/Na/K/Cl/Hct

pH/pCO2/pO2

EasyLyte Accessories		EDMA Code
Catalog No.	Accessory	
2070	EasyLyte EasySampler	21 04 10 01
2101	EasyLyte K+ Electrode	11 04 01 06
2102	EasyLyte Na+ Electrode	11 04 01 07
2113	EasyLyte Cl- Electrode	11 04 01 03
2106	EasyLyte U+ 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
2103	EasyLyte Reference Electrode	11 04 04 01
2258	EasyLyte Membrane Assembly	21 04 10 01
2120	EasyLyte Na/K 800mL Solutions Pack	11 03 01
2121	EasyLyte Na/K/Cl 800mL Solutions Pack	11 03 01
2122	EasyLyte Na/K/Li 800mL Solutions Pack	11 03 01
2123	EasyLyte Na/K/Ca/pH 800mL Solutions Pack	11 03 01
2028	EasyLyte Na/K/Cl/Li 800mL Solutions Pack	11 03 01
2109	EasyLyte Na/K 400mL Solutions Pack	11 03 01
2112	EasyLyte Na/K/Cl 400mL Solutions Pack	11 03 01
2115	EasyLyte Na/K/Li 400mL Solutions Pack	11 03 01
2026	EasyLyte Na/K/Cl/Li 400mL Solutions Pack	11 03 01
2814	EasyLyte Bi-Level Quality Control Kit	11 50 02 04
2815	EasyQC Tri-Level Quality Control Kit	11 50 02 04
2843	EasyLyte Quality Control Sample Cups (60)	21 04 10 01
2118	Daily Cleaning Solution Kit	11 04 10 01
2598	EasyLyte Daily Cleaner Cup	21 04 10 01
2108	EasyLyte Solutions Valve	21 04 10 01
2107	EasyLyte Sample Probe	21 04 10 01
2257	EasyLyte Sample Detector	21 04 10 01
2104	EasyLyte Tubing Kit	21 04 10 01
2100	EasyLyte Calcium Tubing Kit	21 04 10 01
2492	EasyLyte Internal Filling Solution (125mL)	11 04 04 90
2309	EasyLyte Wash Solution (50mL)	11 04 04 90
2111	EasyLyte Urine Diluent (500mL)	11 04 04 90
2577	EasyLyte Standard Solution, Urine (50mL)	11 04 04 90
2323	EasyLyte Probe Wipers (6)	21 04 10 01
2541	EasyLyte Printer Paper (5 rolls)	21 04 10 01



EasyLyte Accessories, continued		EDMA Code
Catalog No.	Accessory	
2595	EasyLyte EasySampler Sample Cups, 500uL (500)	21 04 10 01
2596	EasyLyte Sample Cups 2.0mL (500)	21 04 10 01
10745	Anti-Evaporation Caps (500)	21 04 10 01
2293	EasyLyte Capillary Tubes	21 04 10 01
2590	EasyLyte Capillary Adaptor Kit	21 04 10 01
2292	EasyLyte Capillary Adaptor Cleaning Kit	11 04 04 90
2578	EasyLyte Red Dye Test Solution (50mL)	11 04 04 90
2572	EasyLyte Troubleshooting Kit	21 04 10 01
2571	EasyLyte Troubleshooting Kit (Na/K/Ca/pH and Na/K/Cl/Li)	21 04 10 01
2105	EasyLyte Quarterly Operating Kit	21 04 10 01
2095	EasyLyte Maintenance Kit	21 04 10 01
2076	EasyLyte Sample Tray	21 04 10 01
2074	EasyLyte Sample Cup Retainer Ring	21 04 10 01
7118	Daily Rinse/Cleaning Solution Kit	11 01 01 27
2544	EasyLyte C Series Printer Paper (5 rolls)	21 04 10 01

CERTIFICATO N° 505DM05

CERTIFICATE N° 505DM05

Si certifica che il
this is to certify that

Sistema di Gestione per la Qualità

Quality Management System

messso in atto da

implemented by

NUOVA APTACA S.r.l.

Via Monte Bianco, 4 – IT 20900 MONZA (MB)

nella Sede Operativa di

Operative Unit

Regione Monforte, 30 -- IT 14053 CANELLI (AT)

è conforme alla norma

is in compliance with the standard

UNI CEI EN ISO 13485-2016 (ISO 13485-2016)

per i seguenti Processi
concerning the following kinds of Processes

Gestione della fabbricazione e immissione in commercio di tamponi sterili
per il prelievo di campioni biologici in orifici naturali e in ambito chirurgico.

Progettazione e fabbricazione di dispositivi medici diagnostici per laboratori di analisi.

Commercializzazione di dispositivi medici e diagnostici in vitro.
Management of manufacturing and placing on the market of sterile tampons for sampling of biological specimens in
natural orifice and in surgical field. Design and manufacturing of diagnostic medical devices for analysis laboratories.
Marketing of medical and diagnostic devices in vitro.

Il presente Certificato è soggetto al rispetto delle condizioni stabilite dal Regolamento per la certificazione in vigore applicabili.
This Certificate shall satisfy the requirements established in the Rules for the certification in force applicable.
In caso di discordanza tra le lingue utilizzate nella traduzione del contenuto del presente certificato, fare riferimento alla lingua italiana.
In cases of discrepancy between the languages used in the translation of the content of this certificate, please refer to the Italian language.

L'AMMINISTRATORE DELEGATO
MANAGING DIRECTOR

Roberto Cusolito

Dr. Ing. Roberto Cusolito

Data di Prima Emissione

First Issue Date

2007-10-30

Data di Prima Emissione ITALCERT

First Issue Date ITALCERT

2011-10-30

Data di Rinnovo

Renewal Date

2017-10-30

Data di Delibera

Deliberation Date

2019-01-04

Data di Scadenza

Expiration Date

2020-10-29



ISO 9001

Member degli Accordi di Riconoscimento EA, IAF e ILAC
Signatory of EA, IAF and ILAC Mutual Recognition Agreements

ITALCERT S.r.l. | Viale Sarca, 336 – 20126 Milano (MI) | tel. +39 0266104876 | fax. +39 026610479 | www.italcert.it | italcertsr@legaimail.it

CERTIFICATO N° 5055GQ03

CERTIFICATE N° 5055GQ03

Si certifica che il
this is to certify that

Sistema di Gestione per la Qualità

Quality Management System

impresso in atto da
implemented by

NUOVA APTACA S.r.l.

Via Monte Bianco, 4 - IT 20900 MONZA (MB)

nella Sede Operativa di
Operative Unit

Regione Monforte, 30 - IT 14053 CANELLI (AT)

è conforme alla norma
is in compliance with the standard

UNI-EN ISO 9001-2015 (ISO 9001-2015)

per i seguenti Processi

concerning the following kinds of Processes

Gestione della fabbricazione ed immissione in commercio di tamponi sterili
per il prelievo di campioni biologici in orifici naturali e in ambito chirurgico.
Progettazione e fabbricazione di dispositivi medici diagnostici per laboratori di analisi.
Commercializzazione di dispositivi medici e diagnostici in vitro.

Commercializzazione di articoli da laboratorio

Management of the manufacturing and placing on the market of sterile tampons for sampling of biological specimens
in natural orifice and in surgical field. Design and manufacturing of diagnostic medical devices for laboratories
of analysis. Marketing of medical and diagnostic devices in vitro. Marketing of laboratory articles.

Il presente Certificato è soggetto al rispetto delle condizioni stabilite dal Regolamento per la certificazione in vigore applicabili.

This Certificate shall satisfy the requirements established in the Rules for the certification in force applicable.
In case of discordance between the languages used in the translation of the content of this certificate, please refer to the Italian language.

L'AMMINISTRATORE DELEGATO

MANAGING DIRECTOR

Luigi Cusi

Dr. Ing. Roberto Cusulito

Data di Prima Emissione ITALCERT

First Issue Date ITALCERT

2011-10-30

Data di Rinnovo

Renewal Date

2017-10-30

Data di Scadenza

Expiration Date

2020-10-29

Data di Prima Emissione

First Issue Date

1998-07-23

Settore IAF 14 - 29



SGQ N° 023A PRD N° 122B
RGS N° 023C
ISP N° 075E
Membro degli Accordi di Riconoscimento EA, IAF e ILAC
Signature of EA, IAF and ILAC Mutual Recognition Agreements

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