



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 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).

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).

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).

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

Cécile GOUBAULT,
 Directeur Général Délégué
 Managing Director
 Directora General

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**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 / Codigo GMDN
CHOLESTEROL HDL 2G CALIBRATOR	HDLL-0011 / 0041	DOS-CE-HDLL-CAL	44696
CHOLESTEROL LDL 2G CALIBRATOR	LDLL-0011 / 0041	DOS-CE-LDLL-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	CALJ-0550	DOS-CE-CALJ2	47866
ELITROL I	CONT-0060	DOS-CE-ELIT I	47869
ELITROL II	CONT-0160	DOS-CE-ELIT 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 Standard 100 mg/dL	PRTU-0022	DOS-CE-PRTU100	53482
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

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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).

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 here, 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).

DECLARACION 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).

Sées, le 29 juin 2018

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

ELITech Clinical Systems SAS
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Cécile GOUBAULT,
 Directeur Général Délégué
 Managing Director
 Directora General

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

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
CALCIUM ARSENAZO	CALA-0600/0250	DOS-CE-CALA_R&Std	45789
CALCIUM ENVOY	CALA-0850		
CHLORIDE	CHLO-0600/0250	DOS-CE-CHLO_R&Std	60037
IRON CHROMAZUROL	FECA-0600	DOS-CE-FECA	
IRON ENVOY	FEPE-0850	DOS-CE-FEPE-R&Std	54758
IRON FERRENE	FEPE-0230/0600		
IRON FERROZINE	FEFR-0600	DOS-CE-FEFR_R&Std	
MAGNESIUM CALMAGITE	MAGN-0600	DOS-CE-MAGN_R&Std	
MAGNESIUM ENVOY	MAGX-0850		
MAGNESIUM XTLDIYL	MAGX-0230/0600	DOS-CE-MAGX-R&Std	46795

Vc.



Avantor Performance Materials Poland Sp. z o.o.
 Sowińskiego 11
 44-101 Gliwice
 Tel. 48 32 2392 000

Declaration of conformity

Avantor Performance Materials Poland S.A. who is an established manufacturer of reagents and products for diagnostic in vitro located at:

Sowińskiego 11 Street
 44-101, Gliwice
 Poland

Herewith declares the following:
 Reagents mentioned in attached list are labeled with J.T.Baker label, comply with the In Vitro Diagnostic Medical Devices Directive 98/79/EC and the requirements of ISO 13485 Standard.
 This declaration is the basic for CE marking of the In Vitro Diagnostic Medical Devices.
 The products are not part of List A and List B of Annex II of the IVD Directive 98/79/EC but are subject to self registration.

This declaration is valid for all the IVD medical devices described above and which are placed on the market by ourselves on or after the date hereof and which bear the CE marking

Gliwice, Poland

January 25, 2019

Anna Szuba
 Quality Director

Product	Product number	Pack size
Diluid™ 100 Plus	3961	20 L
Diluid™ 22	2990, 9010PC	10 L
Diluid™ 610	3969	20 L
	3969-00	20 L
Diluid™ Abacus	3430, 9020	20 L
	3430, 9010	10 L
	3430-00	20 L
Diluid™ AC 900	3966	20 L
Diluid™ APR	3476, 9020PC	20 L
Diluid™ Azide free	3957	20 L
Diluid™ III Diff	3963	20 L
	3963, 9010	10 L
	3963-00	20 L
Diluid™ Erma	3459, 9020	20 L
	3459-00	20 L
Diluid™ Mindray	3439, 9020PC	20 L
	3439-00	20 L
Diluid™ NR	3483, 9020PC	20 L
	3483-00	20 L
Diluid™ Ruby	2987, 9020PC	20 L
Diluid™ Sheath 3200-4000	3532, 9020	20 L
Diluid™ ST1600/2000	3976	20 L
Sheath D	3495, 9010PC	10 L
Sheath Fluid 3000/3500	3471, 9020PC	20 L
CN-free Lyse Diff AC 900	3998	5 L
CyMet™ 22 CN Free	2986, 0500PE	500 ml
CyMet™ 3000	3489, 9010PC	10 L
CyMet™ 3200 CN free	3823, 1000	1 L
CyMet™ 3500	3639, 5000PC	5 L
CyMet™ 3500 CN free	3825	5 L
CyMet™ 610 CN free	3970-00	10 L
	3977	5 L
CyMet™ Abacus CN free	3431, 1000	1 L
	3431-00	1 L
CyMet™ APR Basis II	3479, 1000PE	1 L
CyMet™ APR CN free	3417, 0500PE	500 ml
CyMet™ APR EO	3478, 1000PE	1 L
CyMet™ ASA	2950, 2500PE	2.5 L
CyMet™ ASB	2951, 0500PE	500 ml
CyMet™ AS CN free	2952, 9010PC	10 L
CyMet™ BSS3 CN free	2962, 0500PE	500 ml
CyMet™ III Diff	3968	1 L
	3968-00	500 ml
CyMet™ III Diff CN free	3511, 1000	1 L
	3511-00	5 L
CyMet™ Erma	3416-00	500 ml
	3416, 0500	500 ml
	3416, 0500	500 ml
CyMet™ H2O	3653, 1000	1 L
CyMet™ KX CN Free	3425-00	500 ml
	3425, 0500	500 ml
CyMet™ Micro	3852, 1000	500 ml
	3852, 1000	1 L
CyMet™ Micro CN free	3863, 1000	1 L
	3863-00	1 L, micros
	3863-00	1 L, micros
CyMet™ Mindray	3441-00	500 ml
CyMet™ Mindray CN Free	3440, 0500PE	500 ml

NE 041-001-11-01
 Number & KES: 041-001-11-01
 Słowo: 041-001-11-01
 X Wydział Geodezyjny AS
 Słowo: 041-001-11-01
 041-001-11-01

Product	Product number	Pack size
CyMet™ NR III	3484.1000PE	1 L
CyMet™ NR III CN Free	3486-00	1 L
CyMet™ NR V	3486.1000PE	1 L
CyMet™ NR V	3485.1000PE	1 L
CyMet™ Rubly CN Free	2988.5000PC	5 L
CyMet™ ST 1600/2000 CN free	3759.5000	5 L
Leucolysse	3475.5000PC	5 L
Leucolysse Rubly	2989.5000PC	5 L
Blanking Solution 1600/2000	3947	20 L
Detector ergo™	3763	5 L
Detector ergo™	3766	1 L
Detector Ergo™ BS	2970.0900PE	900 ml
ProClean™	3900	5 L
ProClean™	3900-00	5 L
ProClean™ Abacus	3768.1000	1 L micros
ProClean™ CD	3432.5000	5 L
ProClean™ CD	3432.1000PE	1 L
ProClean™ CD	3902.0100PE	100 ml
ProClean™ Extra	3862.5000	5 L
ProClean™ Extra	3862.00	20 L
ProClean™ Extra	3862-00	5 L
ProClean™ Plus	3867-00	1 L micros
Rinse Minderay	3901	1 L micros
8-Parameter Control L/N/H	3442.5000PE	100 ml
8-Parameter Control L/N/H	3427/3428/3429	2.5 ml
8-Parameter Control 4xN	3463/3464/3465	2.5 ml
8-Parameter Control 1xL+4xN+1xH	3747	4 x 2.5 ml
8-Parameter Control extended L/N/H	3751	6 x 2.5 ml
3-Diff Control L/N/H	3633/3634/3635	2.5 ml
3-Diff Control L/N/H	3433/3434/3435	2.5 ml
3-Diff Control L/N/H	3502/3503/3504	2.5 ml
CD-Diff Control L/N/H	3421/3422/3423	4.5 ml
CD-Diff Control 2xL+2xN+2xH	3462/3463/3464	2.5 ml
K-Diff Control L/N/H	3836	3.0 ml
K-Diff Control L/N/H	3456/3456/3457	6 x 3.0 ml
Platelet Control - Extended value	3424	2.5 ml
WBC Reduced RBC L/H	3698/3699	5 x 3.0 ml
XE-Diff Control L/N/H	3737/3732/3733	3.0 ml
Cervix Spray Fixative	3869.1200	4.5 ml
10% v/v Buffered Formaldehyde (4% w/v)	3933.1000	12 x 125 ml
3933.9010	1 L	5 L
3933.9020	10 L	10 L
3933.1000MB	20 L	1000 L
3933.9020PE	20 L	20 L
3933.9010LL	10 L	10 L
3933.9020LL	20 L	20 L
3905.2500PE	2.5 L	2.5 L
3905.5000PE	5 L	5 L
3905.9010PE	10 L	10 L

Product	Product number	Pack size
Eosin-Y Alcoholic	3800.1000PE	1 L
Eosin-Y Alcoholic	3800.2500PE	2.5 L
Giemsa	3856.1000	1 L
Giemsa	3856.2500	2.5 L
Hematoxylin er (Mayer)	3870.1805T	180 L
Hematoxylin er (Mayer)	3870.1000	1 L
Hematoxylin er (Mayer)	3870.2500	2.5 L
Hematoxylin Modified (Harris, Gill II)	3873.1000	1 L
Hematoxylin Modified (Harris, Gill II)	3873.2500	2.5 L
Mey-Grimwald	3855.1000	1 L
Mey-Grimwald	3855.2500	2.5 L
Papanicolaou 2A	3554.1000PE	1 L
Papanicolaou 2A	3554.2500PE	2.5 L
Papanicolaou 2B	3555.1000PE	1 L
Papanicolaou 2B	3555.2500PE	2.5 L
Papanicolaou 3B	3556.1000PE	1 L
Papanicolaou 3B	3556.2500PE	2.5 L
Ultraslit™	3921.0500	500 ml
Ultraslit™	3921.0600	6 x 100 ml
Ultraslit™	3921.9025ST	28 L
Mounting medium High	3882.0500	500 ml
Mounting medium Low	3883.0500	500 ml
PBS	3059	20 L
PBS	3059.9010PC	10 L

Declaration of Conformity



HL-7-0221DC DOI 2015/08 (4)

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
5377	Thrombin Time	55987

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

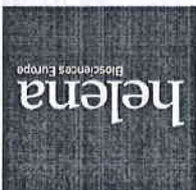
Full Name: M.J. Stephenson Title: Managing Director

Signed:

Date: 10 Aug 2015

Helena Biosciences Europe
Queensway South, Team Valley Trading Estate,
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United Kingdom

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www.helena-biosciences.com



Declaration of Conformity

HL-7-0135 DC DOI 2013/10 (6)

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.

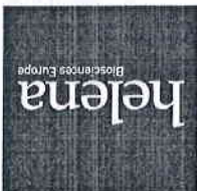
Product Code	Description	GMDN Classification Code
5183	Routine Control SA	30590

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

Date: 31st October 2013

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 United Kingdom



Declaration of Conformity

HL-7-0137 DC DOI 2013/10 (6)

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.

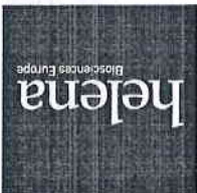
Product Code	Description	GMDN Classification Code
5186	Routine Control N	30590

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: 31st October 2013

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United Kingdom



Declaration of Conformity

HL-7-0138 DC DOI 2013/10 (6)

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: *M.J. Stephenson*

Date: 31st October 2013

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Declaration of Conformity

helena
Biosciences Europe

HL-7-0163 DC DOI 2014/05 (8)

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
5267	Thromboplastin LI	55983
5269	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: 07 May 2014

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United Kingdom

Declaration of Conformity

HL-7-0511 DC DOI 2013/08 (3)

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
5376	Classs Fibrinogen 100	55997
5376H	Classs 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: 05 Aug 2013

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United Kingdom



Declaration of Conformity

HL-7-0700DC DOI 2015/09 (1)



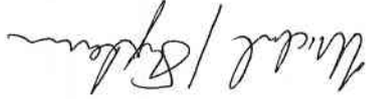
In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.

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
5559SLO	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: 28 Sep 2015

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United Kingdom

EC DECLARATION OF CONFORMITY

Dia.Pro
Diagnostic
 Bio**Pro**bes



MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 - 20099 SESTO SAN GIOVANNI (MILANO) - ITALY
PRODUCT	DIA.BLOOD CODE: DP-9
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, DE MEDICAMENTOS Y PRODUCTOS SANTARIOS)	RELEASED BY AEMPS (AGENCIA ESPAÑOLA
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PLACE & DATE OF FIRST ISSUE	MILANO - NOVEMBER 2010
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) - MARCH 2019
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018

DIA.PRO Diagnostic Bioprobes S.r.l.
 Sede legale e lab.: Via G.Carducci, 27 - 20099 Sesto S.Giovanini (MI) - Italia
 Tel. +39 02 27007161/6450 • Fax +39 02 44386771 • <http://www.diapro.it> • E-mail: info@diapro.it
 Capitale sociale €50.000,00 I.V. - P.IVA: 11924660159 - Reg. Imp. 11924660159 - REA 1509959

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
PROROGA/EXTENSION — Fecha inicial/initial date: 11/12/2003
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

Para el producto/For the product:

Categoría/Category: Productos Sanitarios para Diagnóstico "In Vitro" / In Vitro Diagnostic Medical Devices
Grupo genérico/Genetic 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 Diagnostico Bioprobes S.r.l.
Via G. Carducci, 27 -20099- Sesto San Giovanni – Milano (Italy).

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total Nº 2003 12 0388 CT/ This certificate must be accompanied by the EC Full Quality Assurance System Certificate Nº 2003 12 0388 CT.

Este certificado es consecuencia de la evaluación de la documentación técnica del diseño contenida en el expediente Nº 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 Nº 2003 05 0240, and guarantees that the design of the described products fulfil the requirements of the Directive.

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



Fdo. M^a Jesús Lamas Diaz

Firmado digitalmente por: Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018

Localizador: 62V6ZAG560

CORREO ELECTRONICO

Página 1 de 2

0n0318@semps.es

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel: (+34) 902 101 322 / (+34) 91 822 52 56
Fax: (+34) 91 822 52 56

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
PROROGA/EXTENSION — Fecha inicial/initial date: 11/12/2003
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

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

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

Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis B, mediante técnicas de inmunoadsorción 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]

HBs Ab ELISA cualitativo-cuantitativo / ELISA qualitative-quantitative
- SAB,CE (96 tests)

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

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



Fdo. M^a Jesús Lamas Diaz

Firmado digitalmente por: Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018

Localizador: 62V6ZAG560

CORREO ELECTRONICO

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ORGANISMO NOTIFICADO 0318

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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
PRORROGA/EXTENSION — Fecha inicial/Initial date: 11/12/2003
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

Para el producto/For the product:

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

Elaborado en/In the facilities:

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

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total Nº 2003 12 0388 CT/ This certificate must be accompanied by the EC Full Quality Assurance System Certificate Nº 2003 12 0388 CT.

Este certificado es consecuencia de la evaluación de la documentación técnica del diseño contenida en el expediente Nº 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 Nº 2003 05 0240, and guarantees that the design of the described products fulfil the requirements of the Directive.

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



Fdo. M^o Jesús Lamas Díaz

Firmado digitalmente por Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018
Localizador: B86E8D2586

Puede comprobar la autenticidad del documento en la aplicación Localizador de la Web de la AEMPS
Página 1 de 2
CORREO ELECTRONICO: on0318@aemps.es

ORGANISMO NOTIFICADO 0318

C/ CAMPEZO, 1 - EDIFICIO B
28002 MADRID
Tel: (+34) 902 101 322 / (+34) 91 622 52 59
Fax: (+34) 91 622 52 59

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
PRORROGA/EXTENSION — Fecha inicial/Initial date: 11/12/2003
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

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 infectious diseases

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

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

HCV Ab ELISA cualitativo / ELISA qualitative

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

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

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



Fdo. M^o Jesús Lamas Díaz

Firmado digitalmente por Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018
Localizador: B86E8D2586

Puede comprobar la autenticidad del documento en la aplicación Localizador de la Web de la AEMPS
Página 2 de 2
CORREO ELECTRONICO: on0318@aemps.es

ORGANISMO NOTIFICADO 0318

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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
PRORROGA/EXTENSION — Fecha inicial/Initial date: 04/12/2008
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

Para el producto/For the product:

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

Elaborado en/In the facilities:

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

Este certificado debe ir acompañado por el certificado CE de Sistema de Garantía de Calidad Total Nº 2003-12-0388 CTI. *This certificate must be accompanied by the EC Full Quality Assurance System Certificate Nº 2003-12-0388 CTI.*

Este certificado es consecuencia de la evaluación de la documentación técnica del diseño contenida en el expediente Nº 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 Nº 2003-05-0240, and guarantees that the design of the described products fulfil the requirements of the Directive.*

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



Fdo. M^a Jesús Lamas Díaz

Firmado digitalmente por Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018

Localizador: PELLDBAAS4

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Página 1 de 2

CORREO ELECTRÓNICO
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Fax: (+34) 91 822 52 59
ORGANISMO NOTIFICADO 0318

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
PRORROGA/EXTENSION — Fecha inicial/Initial date: 04/12/2008
Fecha de última prórroga/ Last extension date: 27/11/2013

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

A favor de/In favour of:

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

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

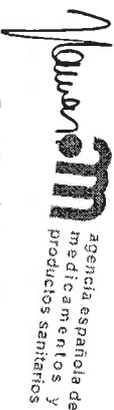
Reactivos y productos reactivos para la determinación, confirmación y cuantificación en muestras humanas de marcadores de infección por Hepatitis B, mediante técnicas de inmunosorción enzimática (EUSAY) Reagents and reactive products for the determination, confirmation and quantification in human specimens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

HBs Ag one Version ULTRA ELISA cualitativo / ELISA qualitative

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

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

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



Fdo. M^a Jesús Lamas Díaz

Firmado digitalmente por Agencia Española de Medicamentos y Productos Sanitarios
Fecha de la firma: 19/11/2018

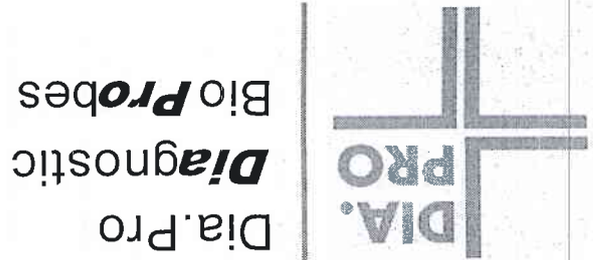
Localizador: PELLDBAAS4

Puede comprobar la autenticidad del documento en la aplicación Localizador de la Web de la AEMPS
Página 2 de 2

CORREO ELECTRÓNICO
01031@emps.es

C/ CAMPEZO, 1 - EDIFICIO 8
28022 MADRID
Tel: (+34) 902 101 322 / (+34) 91 822 52 59
Fax: (+34) 91 822 52 59
ORGANISMO NOTIFICADO 0318

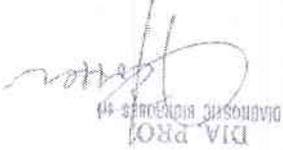
EC DECLARATION OF CONFORMITY



MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 - 20099 SESTO SAN GIOVANNI (MILANO) - ITALY
PRODUCT	HP IgM CODE: HPM.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 - APRIL 2004
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) - MARCH 2019
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018

VECTOR BEST	AO Vector-Best	Rev. 01
EC Declaration of conformity EIA-1-17		Page 1 of 3

EC DECLARATION OF CONFORMITY

AO Vector-Best hereby ensures under own responsibility and declares that the products listed on pages 2-3 are in conformity with applicable provisions and fulfill the essential requirements of Annex I Directive 98/79/EC of 27 October 1998 regarding in vitro diagnostic medical devices.

Classification of products:

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

Harmonized standards applied:

EN ISO 18113-1:2014; EN ISO 18113-2:2011 (In vitro diagnostic medical devices. Information supplied by the manufacturer (labelling). Terms, definitions and general requirements. In vitro diagnostic reagents for professional use); EN ISO 15223-1:2012 (Symbols to be used with medical device labels, labelling and information to be supplied); EN ISO 13485:2012+AC:2012 (Medical devices. Quality management systems. Requirements for regulatory purposes); EN 13612:2002 (Performance evaluation of in vitro diagnostic medical devices); EN 23640:2013 (In vitro diagnostic medical devices. Evaluation of stability of in vitro diagnostic reagents); EN 13641:2002 (Elimination or reduction of risk of infection related to in vitro diagnostic reagents); EN ISO 14971:2012 (Medical devices. Application of risk management to medical devices).

Conformity assessment procedure:

Annex III (not including section 6).

Manufacturer:

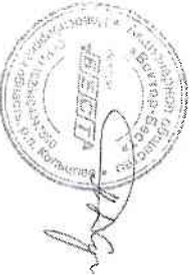
AO Vector-Best
Address: 630559, Koltsovo, Novosibirsk Region, Research and Production area, building 36, office 211, Russian Federation, tel. +7 (383) 336-73-46, tel./fax +7 (383) 332-67-49

European authorized representative:

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

Date: 2017/10/16

Murat Khusainov
General Director AO Vector-Best



Valid until: 2022/07/03

VECTOR BEST	AO Vector-Best	Rev. 01
EC Declaration of conformity EIA-1-17		Page 2 of 3

No.	Product name	Identification data	REF
1.	Veciohep A-IgG	Enzyme immunoassay kit for the qualitative and quantitative determination of IgG to hepatitis A virus	D-0362
2.	VecioMeasles-IgG	Enzyme immunoassay kit for the quantitative and qualitative determination of IgG to measles virus in blood serum (plasma)	D-1356
3.	VecioMeasles-IgM	Enzyme immunoassay kit for the detection of IgM to measles virus in blood serum (plasma)	D-1358
4.	Rotavirus-antigen-EIA-BEST	Enzyme immunoassay kit for the detection of human rotavirus antigen	D-1652
5.	Adenovirus-antigen-EIA-BEST	Enzyme immunoassay kit for the detection of human adenovirus antigen	D-1654
6.	VecioEBV-NA-IgG	Enzyme immunoassay kit for the detection of IgG to nuclear antigen of Epstein-Barr virus in blood serum (plasma)	D-2170
7.	VecioEBV-EA-IgG	Enzyme immunoassay kit for the detection of IgG to early antigen of Epstein-Barr virus in blood serum (plasma)	D-2172
8.	VecioEBV-VCA-IgM	Enzyme immunoassay kit for the detection of IgM to viral capsid antigen of Epstein-Barr virus in blood serum (plasma)	D-2176
9.	VecioMumps-IgG	Enzyme immunoassay kit for the detection of IgG to mumps virus in blood serum (plasma)	D-2602
10.	VecioMumps-IgM	Enzyme immunoassay kit for the detection of IgM to mumps virus in blood serum (plasma)	D-2604
11.	Toxocara-IgG-EIA-BEST	Enzyme immunoassay kit for the detection of IgG to Toxocara antigens in blood serum (plasma)	D-2752
12.	Trichinella-IgG-EIA-BEST	Enzyme immunoassay kit for the detection of IgG to Trichinella antigens in blood serum (plasma)	D-3152
13.	Yersinia-IgG-EIA-BEST	Enzyme immunoassay kit for the detection of IgG to causative agents of yersiniosis	D-3202
14.	Yersinia-IgA-EIA-BEST	Enzyme immunoassay kit for the detection of IgA to causative agents of yersiniosis	D-3204
15.	Yersinia-IgM-EIA-BEST	Enzyme immunoassay kit for the detection of IgM to causative agents of yersiniosis	D-3206
16.	Echinococcus-IgG-EIA-BEST	Enzyme immunoassay kit for the detection of IgG to Echinococcus granulosus antigens in blood serum (plasma)	D-3356
17.	Ascaris-IgG-EIA-BEST	Enzyme immunoassay kit for the detection of IgG to Ascaris lumbricoides antigens in blood serum (plasma)	D-3452
18.	IgA-Transglutaminase-EIA-BEST	Enzyme immunoassay kit for the quantitative determination of IgA to tissue transglutaminase in blood serum (plasma)	D-3758
19.	IgG-Transglutaminase-EIA-BEST	Enzyme immunoassay kit for the quantitative determination of IgG to tissue transglutaminase in blood serum (plasma)	D-3760
20.	Pepsinogen 1-EIA-BEST	Enzyme immunoassay kit for the determination of pepsinogen 1 concentration in blood serum	D-3762
21.	Pepsinogen 2-EIA-BEST	Enzyme immunoassay kit for the determination of pepsinogen 2 concentration in blood serum	D-3764

22.	VectorHanta-IgG	Enzyme immunoassay kit for the detection of IgG to Hantavirus in blood serum (plasma)	D-4902
23.	VectorHanta-IgM	Enzyme immunoassay kit for the detection of IgM to Hantavirus in blood serum (plasma)	D-4904
24.	VectorNile-IgM	Enzyme immunoassay kit for the detection of IgM to West Nile Virus in blood serum (plasma)	D-5150
25.	VectorNile-IgG	Enzyme immunoassay kit for the detection of IgG to West Nile Virus in blood serum (plasma)	D-5152
26.	VectorNile-IgG-avidity	Enzyme immunoassay kit for the determination of avidity index of IgG to West Nile Virus in blood serum (plasma)	D-5154

Certificate

mdc medical device certification GmbH
certifies that

VECTOR



AO Vector-Best
Research and Production area
building 36, Office 211, Koltsovo
630559 Novosibirsk region
Russian Federation

with the locations listed in the attachment
for the scope

Design and development, production and distribution of
medical devices for in vitro diagnostics (PCR, ELISA, Biochemistry)
has introduced and applies a

Quality Management System

The mdc audit has proven that this quality management system
meets all requirements of the following standard

EN ISO 13485

Medical devices – Quality management systems –
Requirements for regulatory purposes

EN ISO 13485:2016 + AC:2015 - ISO 13485:2016

Valid from 2018-02-13
Valid until 2020-07-03
Registration no. D1213100017
Report no. P18-00489-17996
Stuttgart 2018-07-13

Head of Certification Body



mdc medical device certification GmbH
Kraepelstraße 6
D-70191 Stuttgart, Germany
Phone: +49 (0)714 203897-0
Fax: +49 (0)714 203897-10
Internet: <http://www.mdc.de>



Attachment of the certificate
No. D1213100017 date 2018-07-13 Page 1 of 1

Location	Scope
AO Vector-Best, Aduzova str. 1/1, 630117 Novosibirsk, Russian Federation	design and development, production and distribution of medical devices for in vitro diagnostics
AO Vector-Best Research and Production area, building 36, Koltsovo, 630559 Novosibirsk region, Russian Federation	design and development, production of medical devices for in vitro diagnostics
AO Vector-Best, Pasechnaya str. 3, 630117 Novosibirsk, Russian Federation	design and development, production of medical devices for in vitro diagnostics

Head of Certification Body



mdc medical device certification GmbH
Kraepelstraße 6
D-70191 Stuttgart, Germany
Phone: +49 (0)714 203897-0
Fax: +49 (0)714 203897-10
Internet: <http://www.mdc.de>

Сертификат

mdc medical device certification GmbH

Удостоверяет, что на предприятии

ВЕКТОР



АО «Вектор-Бест»

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

с производственными площадками согласно приложению к Сертификату

применительно к областям

проектирование и разработка, производство и реализация

медицинских изделий in-vitro диагностики
(ПЦР, ИФА, биохимия)

была введена и применяется

СИСТЕМА УПРАВЛЕНИЯ КАЧЕСТВОМ

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

EN ISO 13485

Издателя медицинские – Системы менеджмента качества –

Регулирующие системные требования

EN ISO 13485:2016 + АС:2016 – ISO 13485:2016

Дата выдачи	2018-07-13
Срок действия до	2020-07-05
Регистрационный №	D1213100017
Отчет №	P18-00489-117896
Штутгарт, Германия	2018-07-13

Руководитель сертификационного органа



DAKS
Deutscher
Akademischer
Austauschdienst



medical device certification

mdc

mdc medical device certification GmbH
Königsplatz 6

D-73101 Stuttgart, Germany
Phone: +49 (0)71 425552-0
Fax: +49(0)71 425552-10
Internet: <http://www.mdc-cert.de>

№ D1213100017	Приложение к Сертификату от 2018-07-13	Стр. 1 из 1
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Месторасположение	Область действия
АО «Вектор-Бест», ул. Арбузова, 1/1, 630117, г. Новосибирск, Российская Федерация	проектирование и разработка, производство и реализация медицинских изделий in vitro диагностики
АО «Вектор-Бест», 630559, Новосибирская область, р.п. Колыцово, Научно-производственная зона, корпус 36, Российская Федерация	проектирование и разработка, производство медицинских изделий in vitro диагностики
АО «Вектор-Бест», ул. Пасечная, 3, 630117, г. Новосибирск, Российская Федерация	проектирование и разработка, производство медицинских изделий in vitro диагностики



medical device certification

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D-73101 Stuttgart, Germany
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Internet: <http://www.mdc-cert.de>

Руководитель сертификационного органа

Product List – CE Marked

Certified by

ISO 13485:2012

EC – Directive 98 / 79 EC
For In-Vitro-Diagnostics

Prod. No.	Novalisa® Virology Name
ADVA0010	Adenovirus IgA
ADVG0010	Adenovirus IgG
ADVMM0010	Adenovirus IgM
CHIG0590	Chikungunya Virus IgG capture
CHIM0590	Chikungunya Virus IgM µ-capture
CMV/G0110	Cytomegalovirus (CMV) IgG
ACM/V7110	Avidity Cytomegalovirus (CMV) IgG
CMV/M0110	Cytomegalovirus (CMV) IgM
DEN/G0120	Dengue Virus IgG
DEN/M0120	Dengue Virus IgM
DVM0640	Dengue Virus IgM µ-capture
EBVA0150	Epstein-Barr Virus (VCA) IgA
EBVG0150	Epstein-Barr Virus (VCA) IgG
AEBV7150	Avidity Epstein-Barr Virus (VCA) IgG
EBVM0150	Epstein-Barr Virus (VCA) IgM
EBVG0580	Epstein-Barr Virus (EBNA) IgG
HANG0670	Hantavirus IgG
HANM0670	Hantavirus IgM
HSV/G0250	Herpes simplex Virus 1+2 (HSV) IgG
HSV/M0250	Herpes simplex Virus 1+2 (HSV) IgM
HSV1G0500	Herpes simplex Virus 1 (HSV-1) IgG
HSV1M0500	Herpes simplex Virus 1 (HSV-1) IgM
HSV2G0540	Herpes simplex Virus 2 (HSV-2) IgG
HSV2M0540	Herpes simplex Virus 2 (HSV-2) IgM
INFA0290	Influenza Virus A IgA
INFG0290	Influenza Virus A IgG
INFM0290	Influenza Virus A IgM
INFA0300	Influenza Virus B IgA
INFG0300	Influenza Virus B IgG
INFM0300	Influenza Virus B IgM
MEAG0330	Measles Virus IgG
AMEA7330	Avidity Measles Virus IgG
MEAM0330	Measles Virus IgM
MUM/G0340	Mumps Virus IgG
MUM/M0340	Mumps Virus IgM
PAIA0360	Parainfluenza Virus 1,2,3 IgA
PAIG0360	Parainfluenza Virus 1,2,3 IgG
PAR/G0370	Parovirus B 19 IgG
PAR/M0370	Parovirus B 19 IgM
RSVA0380	Respiratory syncytial Virus IgA
RSVG0380	Respiratory syncytial Virus IgG
RSVM0380	Respiratory syncytial Virus IgM
RUBG0400	Rubella Virus IgG
ARUB7400	Avidity Rubella Virus IgG

2018-02

RUBM0400	Rubella Virus IgM μ -capture
TICG0440	TBE / FSME IgG
TICM0440	TBE / FSME IgM
PTICG044	TBE / FSME IgG plus
VZVA0490	Varicella-Zoster Virus (VZV) IgA
VZVG0490	Varicella-Zoster Virus (VZV) IgG
VZVM0490	Varicella-Zoster Virus (VZV) IgM
ZVG0790	Zika Virus IgG capture
ZVM0790	Zika Virus IgM μ -capture

Novalisa® **Bacteriology**

Prod. No.	Name
BOPA0030	Bordetella pertussis IgA
BOPG0030	Bordetella pertussis IgG
BOPM0030	Bordetella pertussis IgM
BPTA0610	Bordetella pertussis toxin (PT) IgA
BPTG0610	Bordetella pertussis toxin (PT) IgG
BORG0040	Borrelia burgdorferi IgG
BORM0040	Borrelia burgdorferi IgM
BRUG0050	Brucella IgG
BRUM0050	Brucella IgM
CHLA0070	Chlamydia trachomatis IgA
CHLG0070	Chlamydia trachomatis IgG
CHLM0070	Chlamydia trachomatis IgM
CHLA0510	Chlamydia pneumoniae IgA
CHLG0510	Chlamydia pneumoniae IgG
CHLM0510	Chlamydia pneumoniae IgM
CORG0090	Corynebacterium diphtheriae toxin IgG
CORG5009	Corynebacterium diphtheriae toxin 5S IgG
PCORRG009	Corynebacterium diphtheriae toxin 5S IgG plus
COX1G0600	Coxiella burnetii (Q-Fever) Phase 1 IgG
COX2G0600	Coxiella burnetii (Q-Fever) Phase 2 IgG
COX2M0600	Coxiella burnetii (Q-Fever) Phase 2 IgM
HELA0220	Helicobacter pylori IgA
HELG0220	Helicobacter pylori IgG
PHIELA022	Helicobacter pylori IgA plus
PHELG022	Helicobacter pylori IgG plus
LEGG0650	Legionella Pneumophila IgG
LEGM0650	Legionella Pneumophila IgM
LEPG0660	Leptospira IgG
LEPM0660	Leptospira IgM
MYCA0350	Mycoplasma pneumoniae IgA
MYCG0350	Mycoplasma pneumoniae IgG
MYCM0350	Mycoplasma pneumoniae IgM

TETG0430	Clostridium tetani toxin IgG
TETG5043	Clostridium tetani toxin 5S IgG
PTTETG043	Clostridium tetani toxin 5S IgG plus

Novalisa® **Parasites**

Prod. No.	Name
CHAG0560	Chagas (Trypanosoma cruzi) IgG
TRYP0570	Chagas
ENTG0140	Entamoeba histolytica IgG
GIA0160S	Giardia lamblia antigen
LEIG0310	Leishmania infantum IgG
MAL0620	Malaria
TOXA0460	Toxoplasma gondii IgA
TOXG0460	Toxoplasma gondii IgG
ATOX7460	Avidity Toxoplasma gondii IgG
TOXM0460	Toxoplasma gondii IgM μ -capture

Novalisa® **Worms**

Prod. No.	Name
ASCG0020	Ascaris lumbricoides IgG
ECHG0130	Echinococcus IgG
FIL0760	Filariasis
SCHG0410	Schistosoma mansoni IgG
SCHM0410	Schistosoma mansoni IgM
STR00690	Strongyloides
TAEG0420	Taenia solium IgG
TOCG0450	Toxocara canis IgG
TRIG0480	Trichinella spiralis IgG

Novalisa® **Fungi**

Prod. No.	Name
ASPG0680	Aspergillus fumigatus IgG
ASPM0680	Aspergillus fumigatus IgM
CANA0060	Candida albicans IgA
CANG0060	Candida albicans IgG
CANM0060	Candida albicans IgM

Novalisa® Hormones

THYROID HORMONES
(ELISAs for the determination of thyroid hormones and antibodies)

Prod. No.	Name
ATG1010	Anti-TG
ATP01020	Anti-TPO
TSH1030	TSH

Novaline

Prod. No.	Name
TRYG2570	Chagas IgG LineBlot

Hormones

STEROID HORMONES
(ELISAs for the determination of steroid hormones in plasma and serum)

Prod. No.	Name
DNOV001	Cortisol
DNOV002	Testosterone
DNOV003	17 Beta-Estradiol
DNOV004	17-OH Progesterone
DNOV005	DHEA-S
DNOV006	Progesterone
DNOV007	Free Estriol
DNOV008	Androstenedione
DNOV009	Free Testosterone
DNOV011	Total Estriol
DNOV012	Aldosterone

STEROID HORMONES IN URINE
(ELISAs for the determination of steroid hormones in urine)

Prod. No.	Name
DNOV010	Urinary Cortisol

STEROID HORMONES IN SALIVA
(ELISAs for the determination of steroid hormones in saliva)

Prod. No.	Name
DSNOV20	Cortisol Saliva
DSNOV21	Testosterone Saliva
DSNOV22	17 beta-Estradiol Saliva

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DSNOV24	DHEA-S Saliva
DSNOV25	Progesterone Saliva
DSNOV26	Estriol Saliva
DSNOV27	Androstenedione Saliva

PROTEIN HORMONES
(ELISAs for the determination of proteins in plasma and serum)

Prod. No.	Name
DNOV030	LH
DNOV031	FSH
DNOV032	Prolactin
DNOV033	AFP
DNOV034	beta HCG

THYROID HORMONES
(ELISAs for the determination of thyroid hormones and antibodies)

Prod. No.	Name
DNOV051	Free T3
DNOV052	Free T4
DNOV053	Total T3
DNOV054	Total T4
DNOV057	Thyroglobulin

DIABETES MONITORING
(ELISAs for the determination of specific analytes in plasma and serum)

Prod. No.	Name
DNOV111	Insulin
DNOV112	C-Peptide

CIRCULATING IMMUNO COMPLEXES
(ELISAs for the determination of specific analytes in plasma and serum)

Prod. No.	Name
DNOV093	CIC-C1q
DNOV094	CIC-C3d
DNOV096	CH-50

TUMOR MARKERS
(ELISAs for the determination of specific analytes in plasma and serum)

Prod. No.	Name
DNOV060	CEA
DNOV061	CA 125
DNOV062	CA 15-3

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DNOV063 CA 19-9

ZVM0790 Zika Virus IgM I-capture

MISCELLANEOUS
(ELISAs for the determination of specific analytes in plasma and serum)

Prod. No.	Name
DNOV100	Ferritin
DNOV101	HGH
DNOV102	IgE

Novalisa® Autoimmune

Autoimmune
(ELISAs for the determination of specific autoimmune antibodies)

Prod. No.	Name
ATG1010	Anti-TG
ATPO1020	Anti-TPO

Rheumatology
(ELISAs for the determination of specific analytes in plasma and serum)

Prod. No.	Name
RFM3010	Rheumatoid Factor IgM

Novalisa® Recombinant Antigens

Prod. No.	Name
BORG0040	Borrelia burgdorferi IgG
BORM0040	Borrelia burgdorferi IgM
CHAG0560	Chagas (Trypanosoma cruzi) IgG
TRYPO570	Chagas
HANG0670	Hantavirus IgG
HANM0670	Hantavirus IgM
HELA0220	Helicobacter pylori IgA
PHELA022	Helicobacter pylori IgA plus
HSV1G0500	Herpes simplex Virus 1 (HSV-1) IgG
HSV1M0500	Herpes simplex Virus 1 (HSV-1) IgM
HSV2G0540	Herpes simplex Virus 2 (HSV-2) IgG
HSV2M0540	Herpes simplex Virus 2 (HSV-2) IgM
MAL0620	Malaria
STR00690	Strongyloides
TREG0470	Treponema pallidum
ZVIG0790	Zika Virus IgG capture

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Novalisa® Quantitative Assays (WHO standardized)

Prod. No.	Name
BPTA0610	Bordetella pertussis toxin (PT) IgA
BPTG0610	Bordetella pertussis toxin (PT) IgG
CORG0090	Corynebacterium diphtheriae toxin IgG
CORG5009	Corynebacterium diphtheriae toxin 5S IgG
PCORG009	Corynebacterium diphtheriae toxin 5S IgG plus
RFM3010	Rheumatoid Factor IgM
RUBG0400	Rubella Virus IgG
TETG0430	Clostridium tetani toxin IgG
TETG5043	Clostridium tetani toxin 5S IgG
PTETG043	Clostridium tetani toxin 5S IgG plus
TOXG0460	Toxoplasma gondii IgG
ATOX7460	Avidity Toxoplasma gondii IgG
TSH1030	TSH

Novalisa® Quantitative Assays

Prod. No.	Name
ATG1010	Anti-TG
ATPO1020	Anti-TPO
BPTA0610	Bordetella pertussis toxin (PT) IgA
BPTG0610	Bordetella pertussis toxin (PT) IgG
CORG0090	Corynebacterium diphtheriae toxin IgG
CORG5009	Corynebacterium diphtheriae toxin 5S IgG
PCORG009	Corynebacterium diphtheriae toxin 5S IgG plus
HELA0220	Helicobacter pylori IgA
HELG0220	Helicobacter pylori IgG
PHELA022	Helicobacter pylori IgA plus
PHELG022	Helicobacter pylori IgG plus
RFM3010	Rheumatoid Factor IgM
RUBG0400	Rubella Virus IgG
ARUB7400	Avidity Rubella Virus IgG
TETG0430	Clostridium tetani toxin IgG
TETG5043	Clostridium tetani 5S toxin IgG
PTETG043	Clostridium tetani toxin 5S IgG plus
TICG0440	TBE / FSME IgG
PTICG044	TBE / FSME IgG plus
TOXG0460	Toxoplasma gondii IgG
ATOX7460	Avidity Toxoplasma gondii IgG
TSH1030	TSH

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Antigen Assays

Prod. No. Name
GIA0160S Giardia lamblia antigen

BORM0040

Borrelia burgdorferi IgM

Novalisa® Igm µ-capture Assays

Prod. No. Name
CHIM0590 Chikungunya Virus IgM µ-capture
DVM0640 Dengue Virus IgM µ-capture
RUBM0400 Rubella Virus IgM µ-capture
TOXM0460 Toxoplasma gondii IgM µ-capture
ZVM0790 Zika Virus IgM µ-capture

Novalisa® Antibody Assays

Prod. No. Name
ASCG0020 Ascaris lumbricoides IgG
CHAG0560 Chagas (Trypanosoma cruzi) IgG
TRYP0570 Chagas
ENTG0140 Entamoeba histolytica IgG
LEIS0310 Leishmania infantum IgG
MAL0620 Malaria
STR00690 Strongyloides
TAEG0420 Taenia solium IgG
TOCG0450 Toxocara canis IgG
TRIG0480 Trichinella spiralis IgG

Novalisa® Avidity Assays

Prod. No. Name
ACMV7110 Avidity Cytomegalovirus (CMV) IgG
AEBV7150 Avidity Epstein-Barr Virus (VCA) IgG
AMEA7330 Avidity Measles Virus IgG
ARUB7400 Avidity Rubella Virus IgG
ATOX7460 Avidity Toxoplasma gondii IgG

Novalisa® Liquor Diagnostic

Prod. No. Name
BORG0040 Borrelia burgdorferi IgG

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19022018-D6

EC DECLARATION OF CONFORMITY

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

Product name	ASO Latex kit
Catalogue number	031100A

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

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

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

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

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



Eddy Velhuis
Technical Director



EC DECLARATION OF CONFORMITY

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

Product name	RF Latex kit
Catalogue number	830100A

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

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

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

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

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



Eddy Velthuis
Technical Director

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 65



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 Veithuis
Technical Director

Lorne Laboratories Limited
Unit 1 Cutbush Park Industrial Estate
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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 Certificate

Directive 98/79/EC Annex IV, excluding Sections 4 and 6
Full Quality Assurance System
In Vitro Diagnostic Medical Devices

Registration No.: HL 60119814 0001

Report No.: 21265422 001

Manufacturer:
Macherey-Nagel GmbH & Co. KG
Neumann-Neander-Str. 6-8
52355 Düren
Deutschland

Products:
Products for self-testing

(see attachment for products and sites included)

Replaces Certificate, Registration No.: HL 60076687 0001

Expiry Date: 2022-05-28

The Notified Body hereby declares that the requirements of Annex IV, excluding section 4 and 6 of the directive 98/79/EC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex IV, section 5 of the aforementioned directive. For placing on the market of List A devices covered by this certificate an EC design-examination certificate according to Annex IV, section 4 and a verification of manufactured products according to section 6 is required.

Notified Body

Effective Date: 2017-05-29

Date: 2017-05-29

Dipl.-Ing. Sven Hoffmann

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg

TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 98/79/EC concerning in vitro diagnostic medical devices with the identification number 0197.



Doc. 1/1, Rev. 0

TÜV Rheinland
LGA Products GmbH
Tillystraße 2, 90431 Nürnberg

Attachment to
Certificate
Registration No.: HL 60119814 0001
Report No.: 21265422 001

Manufacturer:
Macherey-Nagel GmbH & Co. KG
Neumann-Neander-Str. 6-8
52355 Düren
Deutschland

Products for self-testing:
- single and multi-parameter disposable test strips
for urine analysis
- indicator test strips and papers for measurement
of pH in urine

Additional site for warehousing and logistics:

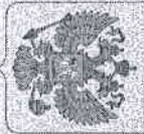
Bahnstr. 120
52355 Düren, Germany

Notified Body

Date: 2017-05-29

Dipl.-Ing. Sven Hoffmann





ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 11 января 2017 года № ФСР 2010/08997

На медицинское изделие
Набор контрольных растворов белков мочи "БМ-контроль"
по ТУ 9398-269-52208224-2010

Настоящее регистрационное удостоверение выдано
Обществу с ограниченной ответственностью "Медякор С-П"
(ООО "Медякор С-П"), Россия,
194100, Санкт-Петербург, ул. А. Матросова, д. 4, корп. 2, Лит. П, офис 212

Производитель
Общество с ограниченной ответственностью "Медякор С-П"
(ООО "Медякор С-П"), Россия,
194100, Санкт-Петербург, ул. А. Матросова, д. 4, корп. 2, Лит. П, офис 212

Место производства медицинского изделия
ООО "Медякор С-П", Россия, 194100, Санкт-Петербург, ул. А. Матросова, д. 4,
корп. 2, Лит. П

Номер регистрационного досье № РД-14955/64156 от 20.12.2016

Вид медицинского изделия 206630

Класс потенциального риска применения медицинского изделия I

Код Общероссийского классификатора продукции для медицинского изделия 93 9816

Настоящее регистрационное удостоверение имеет приложение на 1 листе

Приказом Росздравнадзора от 11 января 2017 года № 80
допущено к обращению на территории Российской Федерации
Заместитель руководителя Федеральной службы
по надзору в сфере здравоохранения



Д.Ю. Павлюков
0024833

ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

ПРИЛОЖЕНИЕ К РЕГИСТРАЦИОННОМУ УДОСТОВЕРЕНИЮ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 11 января 2017 года № ФСР 2010/08997

Лист 1

На медицинское изделие
Набор контрольных растворов белков мочи "БМ-контроль"
по ТУ 9398-269-52208224-2010:

- комплект 1 «БМ-контроль-ССК»;
- комплект 2 «БМ-контроль-ССК + глюкоза и рН»;
- комплект 3 «БМ-контроль-ССК с калibratorом»;
- комплект 4 «БМ-контроль-ССК + глюкоза и рН с калibratorом»;
- комплект 5 «БМ-контроль-ЛПК»;
- комплект 6 «БМ-контроль-ЛПК + глюкоза и рН».

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Заместитель руководителя Федеральной службы
по надзору в сфере здравоохранения



Д.Ю. Павлюков
0026953



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВОХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 15 сентября 2016 года № РЗН 2016/4742

На медицинское изделие

Станки и колбы стеклянные лабораторные по ТУ 9464-019-29508133-2015

Настоящее регистрационное удостоверение выдано

Обществу с ограниченной ответственностью "МиниМед"

(ООО "МиниМед"), Россия,

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

Прозодителю

Общество с ограниченной ответственностью "МиниМед"

(ООО "МиниМед"), Россия,

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

Место производства медицинского изделия

см. приложение

Номер регистрационного досье № РД-7577/28289 от 25.06.2015

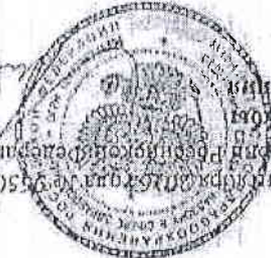
Вид медицинского изделия -

Класс потенциального риска применения медицинского изделия I

Код Общероссийского классификатора продукции для медицинского изделия 94 6456

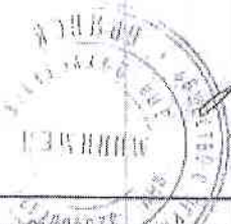
Настоящее регистрационное удостоверение имеет приложение на 1 листе

приказом Росздравнадзора от 15 сентября 2016 года № 2450
покупено к обращению на территории Брянской области.
Руководитель Федеральной службы
по надзору в сфере здравоохранения



М.А. Мурашко

0020781





ФЕДЕРАЛЬНОЕ АГЕНТСТВО
ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ

СВИДЕТЕЛЬСТВО

об утверждении типа средства измерений

RU.C.29:000.A № 30182

Срок действия до 02 октября 2017 г.

НАИМЕНОВАНИЕ ТИПА СРЕДСТВ ИЗМЕРЕНИЙ
Цилиндры исполнений 1, 3

ИЗГОТОВИТЕЛЬ

Общество с ограниченной ответственностью "МиниМедПром"
(ООО "МиниМедПром"), г. Дятьково, Брянская область

РЕГИСТРАЦИОННЫЙ № 24176-07

ДОКУМЕНТ НА ПОВЕРКУ
ГОСТ 8.234-77

ИНТЕРВАЛ МЕЖДУ ПОВЕРКАМИ
Периодическая поверка до ввода в эксплуатацию

Тип средств измерений утвержден приказом Федерального агентства по
техническому регулированию и метрологии от 02 октября 2012 г. № 823

Описание типа средств измерений является обязательным приложением
к настоящему свидетельству.

Заместитель Руководителя
Федерального агентства

Ф.В. Булыгин

"12" 2012 г.

Серия СИ



006811



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВОХРАНЕНИЯ
(РОСЗНАРНАДЗОР)

**РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ
НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ**

№ РЗН 2013/1034

от 12 августа 2013 года

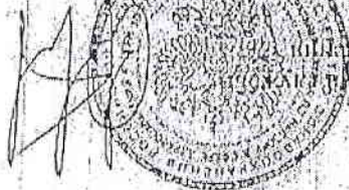
Историческое предприятие зарегистрировано в виде
Открытого акционерного общества «Стр. Лансе», Россия,
242640, Брянская область, Дятьковский район, пос. Стрп, ул. Ленин, д. 6
и производит, чья медицинское изделие
Число Регистрации ЧДН-2 100*20 по ГОСТ 23932-90
Историческая

242640, Брянская область, Дятьковский район, пос. Стрп, ул. Ленин, д. 6
242640, Брянская область, Дятьковский район, пос. Стрп, ул. Ленин, д. 6
242640, Брянская область, Дятьковский район, пос. Стрп, ул. Ленин, д. 6
является производителем изделия

ОКН 94 6450

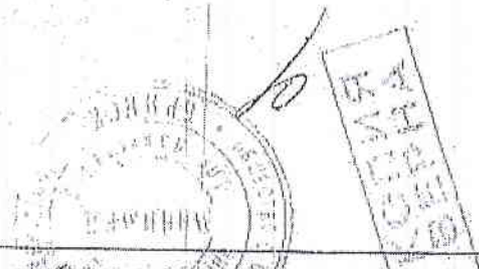
для медицинского изделия -

содержащее регистрационный номер № 40945 от 19.11.2012
приказа Федеральной службы по надзору в сфере здравоохранения
от 12 августа 2013 года № 3990-П/13



А.В. Ларохинко

0002396





ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ

от 13 октября 2010 года № ФСР 2010/09012

На медицинское изделие
Счетчики лабораторные С-5, С-5М по ТУ 9443-005-39766267-2010

Настоящее регистрационное удостоверение выдано

Общество с ограниченной ответственностью "Стилум Плюс"

(ООО "Стилум Плюс"),

Россия, 142290, Московская область, г. Пушкино, ул. Институтская, д. 7

Производитель

Общество с ограниченной ответственностью "Стилум Плюс"

(ООО "Стилум Плюс"),

Россия, 142290, Московская область, г. Пушкино, ул. Институтская, д. 7

Место производства медицинского изделия

ООО "Стилум Плюс", 142290, Московская область, г. Пушкино,

ул. Институтская, д. 7

Номер регистрационного досье № 55566 от 24.08.2010

Вид медицинского изделия -

Класс потенциального риска примененная медицинское изделие 2а

Код Общероссийского классификатора продукции для медицинского изделия 94 4330

приказом Росздравнадзора от 13 октября 2010 года № 9805-1П/10
и приказом от 15 сентября 2016 года № 9678 о замене
допущено к обращению на территории Российской Федерации.

Руководитель Федеральной службы
по надзору в сфере здравоохранения

М.А. Мурашко

0028270

0026420

Д.Ю. Павлюков

приказом Росздравнадзора от 13 октября 2011 года № 6599-ПР/11
и приказом от 05 декабря 2016 года № 13749 о замене
допущено к обращению на территории Российской Федерации
Заместитель руководителя Федеральной службы
по надзору в сфере здравоохранения

Настоящее регистрационное удостоверение имеет приложение на 1 листе

Код Общероссийского классификатора продукции для машинного изделия 94 3790

Класс потенциального риска применения машинного изделия I

Вид машинного изделия -

Номер регистрационного досье № 12259 от 11.04.2011

Место производства машинного изделия
ООО "БАКТЕР", Россия, 248000, г. Калуга, проезд 2-й Академический, д. 17

Производитель
Общество с ограниченной ответственностью "БАКТЕР" (ООО "БАКТЕР"),
Россия, 248000, г. Калуга, проезд 2-й Академический, д. 17

Общество с ограниченной ответственностью "БАКТЕР" (ООО "БАКТЕР"),
Россия, 248000, г. Калуга, проезд 2-й Академический, д. 17

Настоящее регистрационное удостоверение выдано
"Бактер" в комплектах и в отдельных упаковках по ТУ 9437-001-82867591-2010
Изделия машинные для взятия биопров на бактериологические исследования
На машинное изделие

РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ от 13 октября 2011 года № ФСР 2011/12125

ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)



ФЕДЕРАЛЬНАЯ СЛУЖБА ПО НАДЗОРУ В СФЕРЕ ЗАРОВООХРАНЕНИЯ
(РОСЗДРАВНАДЗОР)



РЕГИСТРАЦИОННОЕ УДОСТОВЕРЕНИЕ НА МЕДИЦИНСКОЕ ИЗДЕЛИЕ от 28 октября 2013 года № ФСР 2009/04472

На медицинское изделие

Набор реактивов для титровой пробы "Титровая проба-Агат"

Настоящее регистрационное удостоверение выдано

Обществу с ограниченной ответственностью "Агат-Мед"

(ООО "Агат-Мед"), Россия,

105173, г. Москва, поселок Восточный, ул. Лавная, д. 6, кв. 12

Производитель

Общество с ограниченной ответственностью "Агат-Мед"

(ООО "Агат-Мед"), Россия,

105173, г. Москва, поселок Восточный, ул. Лавная, д. 6, кв. 12

Место производства медицинского изделия

105173, г. Москва, поселок Восточный, ул. Лавная, д. 6, кв. 12

Номер регистрационного Lose № РЛ-2080/39533 от 23.10.2013

Вид медицинского изделия -

Класс потенциального риска применения медицинского изделия 2а

Код Общероссийского классификатора продукции для медицинского изделия 93 9816

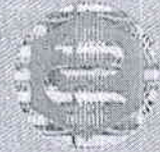
приказом Росздравнадзора от 28 октября 2013 года № 6179-ПР/13
допущено к обращению на территории Российской Федерации.



M.A. Muryashko

M.A. Muryashko
0005084

Врио руководителя Федерального агентства
по надзору в сфере здравоохранения



№ 000548

СЕРТИФІКАТ ВІДПОВІДНОСТІ
Certificate of Conformity

№ UA.TR.101-388.489-2019

Дата реєстрації 11.06.2019 р.

Термін дії до 10.06.2024 р.

Продукція
*Production***Вироби для самоконтролю для діагностики in vitro:**
Смужки індикаторні «Глюкотест», Смужки індикаторні «Ацетонтест»,
Смужки індикаторні «Прототест», Смужки індикаторні «рН-тест»,
Смужки індикаторні «Денситест», Смужки індикаторні «Лаборапт»; Смуги
індикаторні «ТЕМОГЛАН», Набори для визначення йоду у сечі «ЙОДТЕСТ»**Відповідає вимогам**
*Comply with the requirements*Технічного регламенту щодо медичних виробів для діагностики in vitro,
затвердженого Постановою КМУ від 2 жовтня 2013 р. № 754**Виробник**
*Producer(s)*Товариства з обмеженою відповідальністю «ПВП «НОРМА»
(ТОВ «НОРМА»)
вул. Котельникова, буд. 46, м. Київ, 03115, Україна**Місце виробництва**
*Place of production*Товариства з обмеженою відповідальністю «ПВП «НОРМА»
(ТОВ «НОРМА»)
проспект Леся Курбаса, 2Б, м. Київ, 03148, Україна. ЄДРПОУ 16292890.**Сертифікат видано**
органом з оцінки
відповідності
Certificate is issued by the
*conformity assessment body*Державним українським об'єднанням «Політехмед»
(«ДУО «Політехмед»)**На підставі**
*On the grounds of*Оцінки та сертифікації комплексної системи управління якістю згідно з Додатком 4
до Технічного регламенту щодо медичних виробів для діагностики in vitro. Рішення
щодо надання сертифікації від 11.06.2019 р.Нагляд за сертифікованою системою управління якістю здійснюється з
періодичністю, яка регламентується програмою нагляду**Р. Карпавчен****Генеральний директор**
ДУО «Політехмед»**Керівник Органу з оцінки відповідності**

SGQ N° 004A

MEMBRO DEGLI ACCORDI DI MUTUA RICONOSCIMENTO EA, IAF E ILAC
SIGNATORY OF EA, IAF AND ILAC MUTUAL RECOGNITION AGREEMENTS



Data emissione
18/01/2007

Emissione corrente
18/01/2019

Data di scadenza
17/01/2022

ICIM S.p.A.
Piazza Don Enrico Mapelli, 75 - 20099 Sesto San Giovanni (MI)
www.icim.it

Riferirsi alla documentazione del Sistema di Gestione per la Qualità aziendale per l'applicabilità dei requisiti della norma di riferimento.
Refer to the documentation of the Quality Management System for details of application to reference standard requirements.
Per informazioni puntuali e aggiornate circa eventuali variazioni intervenute nello stato della certificazione di cui al presente certificato, si prega di contattare il n° telefonico +39 02 725341 o indirizzo e-mail info@icim.it.
For timely and updated information about any changes in the certification status referred to in this certificate, please contact the number +39 02 725341 or email address info@icim.it.

Progettazione e produzione di provette con vuoto prelievo ematico, liquidi biologici e urine. Produzione di provette per microprelievi di sangue. Commercializzazione di prodotti del Gruppo: kit diagnostici, terreni di coltura per microbiologia, articoli in plastica per laboratorio analisi, provette con vuoto predefinito e aghi sterili.
Design and production of test tubes with predefinit vacuum for collection of haematological samples, biological liquids and urine samples. Trading of the products of the Group: diagnostic kits, culture media for haematology, plastic disposable labware, test tubes with predefinit vacuum and sterile needles.

EA: 14 - 29

PER LE SEGUENTI ATTIVITÀ / FOR THE FOLLOWING ACTIVITIES
Sistema di Gestione per la Qualità / Quality Management System

UNI CEI EN ISO 13485:2016

È CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

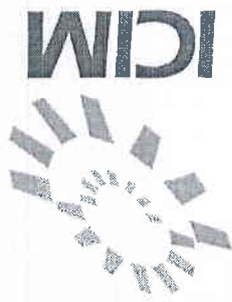
Sede / Head office
Via dell'Industria, 12 - 35020 Arzegrande (PD) - Italia
Uffici direzionali e amministrativi
Unità Operative / Operative Units
Via dell'Industria, 12 - 35020 Arzegrande (PD) - Italia
Progettazione e produzione di provette con vuoto prelievo ematico, liquidi biologici e urine. Produzione di provette per microprelievi di sangue. Commercializzazione di prodotti del Gruppo: kit diagnostici, terreni di coltura per microbiologia, articoli in plastica per laboratorio analisi, provette con vuoto predefinito e aghi sterili.
Via Leonardo Da Vinci, 22 - 35028 Piove di Sacco (PD)
Uffici commerciali e magazzino.

VACUTEST KIMA S.r.l.

SI CERTIFICA CHE IL SISTEMA DI GESTIONE PER LA QUALITÀ DI
WE HEREBY CERTIFY THAT THE QUALITY MANAGEMENT SYSTEM OPERATED BY

CERTIFICATO N.
CERTIFICATE NO.

4265/4/D



CISQ is a member of

IQNet, the association of the world's first class certification bodies, is the largest provider of management System Certification in the world. IQNet is composed of more than 30 bodies and counts over 150 subsidiaries all over the globe.

www.iqnet-certification.com

CISQ è la Federazione Italiana di Organismi di Certificazione dei sistemi di gestione aziendale. CISQ is the Italian Federation of management System Certification Bodies.



www.cisq.com



CISQ is a member of



IQNet, the association of the world's first class certification bodies, is the largest provider of management System Certification in the world. IQNet is composed of more than 30 bodies and counts over 150 subsidiaries all over the globe.

CERTIFICATO n.
CERTIFICATE No.

4264/4

SI CERTIFICA CHE IL SISTEMA DI GESTIONE PER LA QUALITÀ DI
WE HEREBY CERTIFY THAT THE QUALITY MANAGEMENT SYSTEM OPERATED BY

GRUPPO VACUTEST KIMA

Sede / Head Office

Via dell'Industria, 12 – 35020 Arzergrande (PD) - Italia

Unità Operative / Operative Units

MEUS S.r.l. - Via Leonardo da Vinci, 24B – 26 – 28 – Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia

MEUS S.r.l. - Via dell'Industria, 2 - 16 – 35020 Arzergrande (PD) - Italia

ROLL S.R.L. - Via Leonardo Da Vinci, 24A – Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia

KIMA S.R.L. - Via Leonardo da Vinci, 22 – Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia

VACUTEST KIMA S.r.l. - Via dell'Industria, 12 – 35020 Arzergrande (PD) – Italia

VACUTEST KIMA S.r.l. via L. Da Vinci, 22 Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia

È CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

UNI EN ISO 9001:2015

Sistema di Gestione per la Qualità / *Quality Management System*

PER LE SEGUENTI ATTIVITÀ / *FOR THE FOLLOWING ACTIVITIES*

EA: 14 - 29

Progettazione e produzione di provette con vuoto predeterminato ad uso prelievo ematico, liquidi biologici e urine. Progettazione di provette per microprelievi di sangue. Progettazione e produzione di Holders (camicie) per prelievo sottovuoto. Progettazione e produzione di kit diagnostici per l'analisi del sangue e dei liquidi biologici. Progettazione e produzione di terreni di coltura per microbiologia. Progettazione e produzione di aghi e dispositivi sterili per il prelievo ematico. Commercializzazione di prodotti del Gruppo: kit diagnostici, terreni di coltura per microbiologia, articoli in plastica per laboratorio analisi, provette con vuoto predeterminato e aghi sterili. Progettazione e produzione di stampi per articoli in plastica per laboratorio analisi. Stampaggio di materie termoplastiche ad iniezione per articoli medicali.

Design and production of test tubes with predetermined vacuum for collection of haematological samples, biological liquids and urine samples. Production of test tubes for micro-collection of haematological samples. Design and production of Holders for vacuum sampling. Design and production of diagnostic kits for blood and biological liquids analysis. Design and production of culture media for microbiology. Design and production of sterile needles and devices for collection of haematological samples. Trading of the products of the Group: diagnostic kits, culture media for microbiology, plastic disposable labware, test tubes with predetermined vacuum and sterile needles. Design and production of moulds for plastic labware. Injection moulding of thermoplastic materials for medical devices.

Riferirsi alla documentazione del Sistema di Gestione per la Qualità aziendale per l'applicabilità dei requisiti della norma di riferimento.
Refer to the documentation of the Quality Management System for details of application to reference standard requirements.

Il presente certificato è soggetto al rispetto del documento ICIM "Regolamento per la certificazione dei sistemi di gestione" e al relativo Schema specifico.
The use and the validity of this certificate shall satisfy the requirements of the ICIM document "Rules for the certification of company management systems" and specific Scheme.

Per informazioni puntuali e aggiornate circa eventuali variazioni intervenute nello stato della certificazione di cui al presente certificato, si prega di contattare il n° telefonico +39 02 725341 o indirizzo e-mail info@icim.it

For timely and updated information about any changes in the certification status referred to in this certificate, please contact the number +39 02 725341 or email address info@icim.it.

Data emissione
First issue
18/01/2007

Emissione corrente
Current issue
18/01/2019

Data di scadenza
Expiring date
17/01/2022

ICIM S.p.A.

Piazza Don Enrico Mapelli, 75 – 20099 Sesto San Giovanni (MI)
www.icim.it



SGQ N° 004 A

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



www.cisq.com

CISQ è la Federazione Italiana di Organismi di Certificazione dei sistemi di gestione aziendale.
CISQ is the Italian Federation of management system Certification Bodies.

CERTIFICATO N° 505SGQ03

CERTIFICATE N° 505SGQ03

Si certifica che il
this is to certify that

Sistema di Gestione per la Qualità

Quality Management System

messo 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 orifizi naturali e in ambito chirurgico.
Progettazione e fabbricazione di dispositivi medico diagnostici per laboratori di analisi.
Commercializzazione di dispositivi medici e diagnostici in vitro.

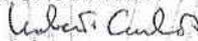
Commercializzazione di articoli da laboratorio

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 dai Regolamenti 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



Dr. Ing. Roberto Cusolito

Data di Prima Emissione
First Issue Date

1998-07-23

Settore IAF 14 - 29

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



SGQ N° 023A PRD N° 122B
SGA N° 020D ISP N° 075E
PAS N° 097C

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



CERTIFICATO N° 505DM05

CERTIFICATE N° 505DM05

Si certifica che il
this is to certify that

Sistema di Gestione per la Qualità

Quality Management System

messo 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 orifizi naturali e in ambito chirurgico.

Progettazione e fabbricazione di dispositivi medico 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 dai Regolamenti 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

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



SGQ N° 023A

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

SGS

Certificado ES10/81672

The management system of

DELTALAB, S.L.

Pol. Ind. La Llana, Plaza De La Verueda, 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. Commercialization of equipment for the storage of prepared samples, cryogenic stored samples, general labware and industrial packages.

Diseño, fabricación y comercialización de material de laboratorio para la toma, transporte y conservación de muestras para análisis de microbiología, biología molecular, hematología, bioquímica, histología, microscopía y coloración. Comercialización de equipos para el almacenamiento de muestras preparadas, almacenamiento de muestras para criogenización, material general de laboratorio y envases industriales.
in/ from the following sites

Pol. Ind. La Llana, Plaza De La Verueda 1 - 08191 Rubí (Barcelona)

This certificate is valid from
29 November 2017 until 11 October 2019.
Issue 7. Certified since October 2010.

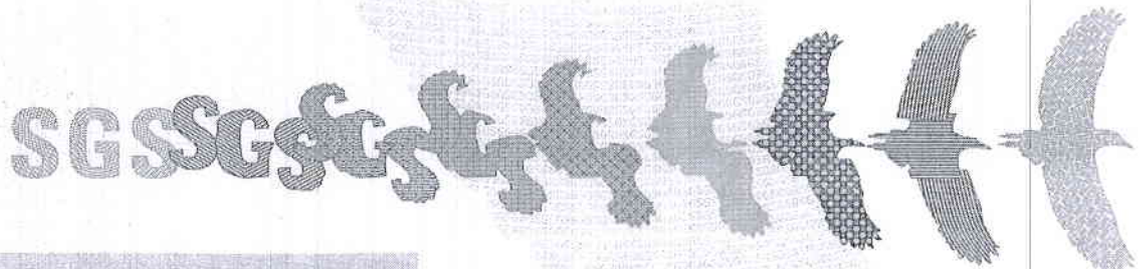
Este certificado es válido desde
29 de noviembre de 2017 hasta 11 de octubre de 2019.
Edición 7. Certificado desde octubre de 2010..

Authorized by

Dirección de Certificación

SGS ICS Ibérica, S.A. (Unipersonal)
C/Trespademe, 29. 28042 Madrid. España.
t 34 91 313 8115 f 34 91 313 8102 www.sgs.com

Page 1 of 1



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SGS

Certificate ES10/81671

The management system of

DELTALAB, S.L.

Polígono Industrial La Llana, Plaza De La Veneda 1,
08191 Rubí, Barcelona. Spain

has been assessed and certified as meeting the requirements of

ISO 13485:2016 EN ISO 13485:2016



For the following activities

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.

Diseño, fabricación y comercialización de productos sanitarios estériles y no estériles para la toma, transporte y conservación de muestras biológicas para análisis clínicos y de IVD.

This certificate is valid from 18 September 2017 until 11 October 2019
and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 10 September 2019

Issue 7. Certified since 12 October 2010

Authorised by

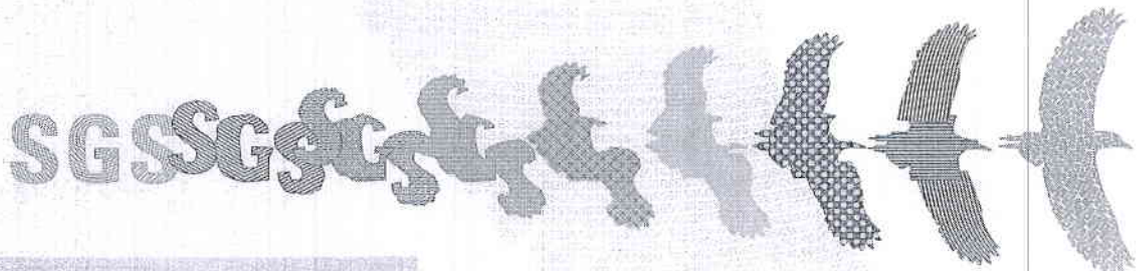


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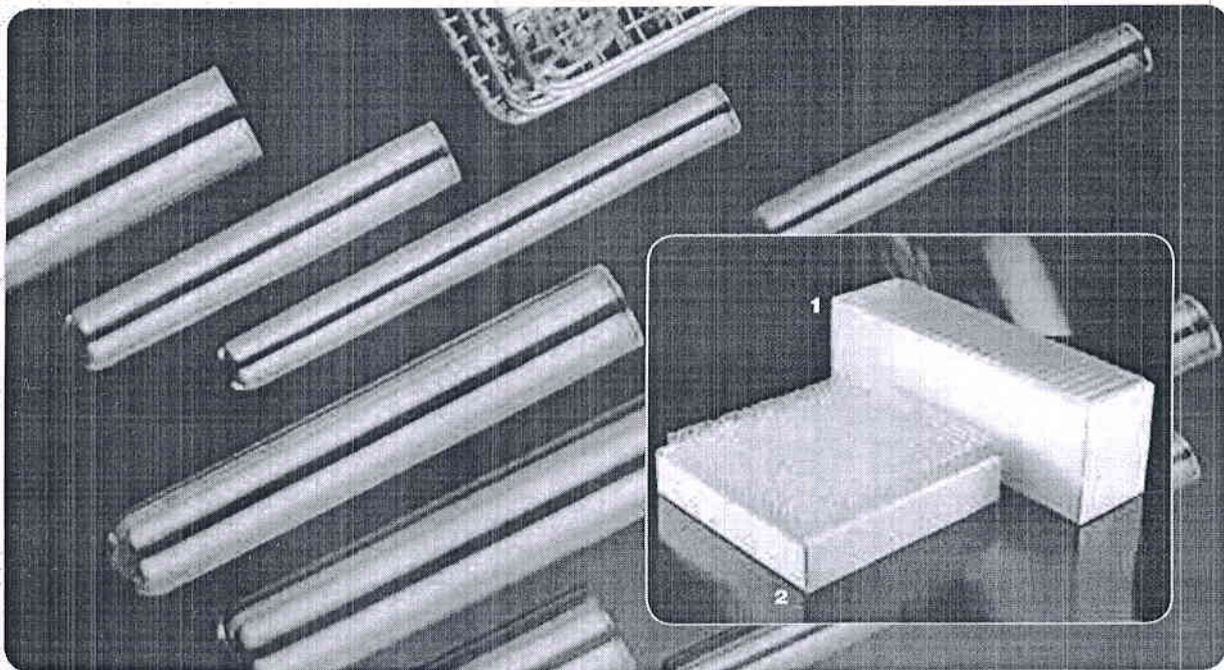
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SGS 13485 2016 0417

Page 1 of 1



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Round bottom glass tubes

Made of **borosilicate** or **soda** glass.

The high quality of those tubes is reflected in the uniformity of their wall thickness and of their diameter and height dimensions.

Supplied in small quantities per case for a more convenient use in laboratory.



code soda	total capacity ml	Ø int. mm tube	Ø ext. mm tube	height mm	thickness mm	case quantity	case weight	case volume
Supplied in boxes (1)								
801075	4	8.20	9.75	75	0.60	4 x 250	3.60	0.010
801275	6	10.20	11.60	75	0.60	4 x 250	4.50	0.013
813100	10	11.10	12.70	100	0.60	4 x 250	6.59	0.022
816100	15	13.95	15.75	100	0.60	4 x 250	9.10	0.034
816150	22	13.55	16.00	150	0.70	4 x 250	13.60	0.049
816160	27	14.40	16.00	160	0.55	500	5.50	0.018
818150	28	15.00	18.00	150	0.85	2 x 250	7.30	0.030
820150	34	17.20	20.00	150	0.85	100	1.92	0.006
820200	47	17.15	19.25	200	0.85	250	6.30	0.020
Supplied in trays (2)								
801175T	6	10.10	11.60	75	0.50	550	1.89	0.005

code boro	total capacity ml	Ø int. mm tube	Ø ext. mm tube	height mm	thickness mm	case quantity	case weight	case volume
Supplied in boxes (1)								
901075	4	8.20	9.75	75	0.60	4 x 250	3.60	0.010
901275	6	10.20	11.60	75	0.60	4 x 250	4.50	0.013
913100	10	11.10	12.70	100	0.60	4 x 250	6.59	0.022
916100	15	13.95	15.75	100	0.60	4 x 250	9.10	0.034
916150	22	13.55	16.00	150	0.70	4 x 250	13.60	0.049
918150	28	15.00	18.00	150	0.85	4 x 125	7.30	0.040

1. Назначение

Коробки стерилизационные круглые КСК-3, КСК-6, КСК-9, КСК-12, КСК-18 (далее – коробки) предназначены для размещения в них перевязочного материала, операционного белья, хирургического инструмента и других предметов медцинского назначения с целью стерилизации их в паровых медцинских стерилизаторах и доставки к месту использования, а также стерильного хранения в течение 3 суток.

Регистрационное удостоверение МЗ РБ № ИМ-7.5619/1004 от 30.03.2010.

Свидетельство о государственной регистрации МЗ Украины № 6184/2007 от 13.03.2012 г.

Регистрационное удостоверение МЗ Казахстана РК-ИМН-5-№ 001180 от 04.02.2013 г.

Регистрационное удостоверение МЗ Узбекистана № ТГ 13507 от 04.06.2012.

2. Технические характеристики

Наименование параметра	КСК-3	КСК-6	КСК-9	КСК-12	КСК-18
Условный объем, дм ³	3	6	9	12	18
Диаметр, мм	169 _s	234 _s	269 _s	319 _s	374 _s
Масса, кг	0,7±0,1	1,1±0,1	1,3±0,1	1,7±0,15	2,35±0,15

3. Комплект поставки

Коробка, шт.
Пасторт, шт.
Упаковка, шт.

1
1
1

4. Подготовка коробок к работе

До начала применения, коробки должны подвергаться химической очистке дезинфекцией и предстерилизационной обработке в соответствии с действующими инструкциями Минздрава.

5. Свидетельство о приемке, упаковке и продаже

Коробка стерилизационная круглая КСК - 9
соответствует требованиям ТУ 92-0024346-04-93 «Коробки стерилизационные круглые КСК» и предназначена годной к эксплуатации.

Печать ОТК

Дата упаковки (готовления)

Продана

(наименование предприятия торговли)

Дата продажи

(печать магазина)

(подпись)

Общий срок хранения коробок не ограничен.

6. Гарантия изготовителя

6.1 Гарантийный срок – 24 месяца с даты продажи коробок, но не более 36 месяцев с даты изготовления.

6.2 Гарантийный срок хранения – 24 месяца с даты продажи коробок.

6.3 В течение гарантийного срока изготовитель ремонтирует или заменяет коробки, или их составные части, пришедшие в негодность по вине изготовителя, при предъявлении настоящего паспорта.

Liofilchem®

DICHIARAZIONE DI CONFORMITÀ CE / EC DECLARATION OF CONFORMITY

DICHIARAZIONE DI CONFORMITÀ CE

La società Liofilchem® S.r.l., con Sede Legale in Via Scozia, 64026 Roseto degli Abruzzi (TE) Italia, in qualità di fabbricante del dispositivo medico-diagnostico *in vitro* elencato nella tabella allegata Revisione 31.0 del 08.01.2016

dichiara sotto la propria responsabilità

- che il dispositivo sopra indicato soddisfa tutte le disposizioni applicabili della Direttiva 98/79/CE (Allegato II) recepita nella Legislazione Italiana dal Decreto Legislativo n° 332 del 8 settembre 2000;
- che il dispositivo in oggetto non è incluso nell'Allegato II, lista A e B della Direttiva 98/79/CE
- che la documentazione tecnica di cui all'allegato III della direttiva Direttiva 98/79/CE è a disposizione delle autorità nazionali presso la sua sede e sarà conservata per 5 anni dall'ultima data di fabbricazione del prodotto;
- che il processo di fabbricazione segue adeguati principi di assicurazione della qualità;
- di aver attuato e di mantenere aggiornato, un sistema di sorveglianza post-produzione per il monitoraggio dei prodotti;
- che il dispositivo in oggetto è stato messo in commercio munito di marcatura CE.

EC DECLARATION OF CONFORMITY

The company Liofilchem® S.r.l., registered office in Via Scozia, 64026 Roseto degli Abruzzi (TE) Italy, as a manufacturer of the *in vitro* medical diagnostic device listed in the attached table, Revision 31.0 of 08.01.2016

hereby certifies under its own responsibility

- that the above mentioned device complies with all the applicable provisions of Directive 98/79/EC (Annex III) and its relevant transposition into national law;
- the above mentioned is not included in Annex II, List A and B of Directive 98/79/EC;
- that the technical documentation referred to at Annex III of the Directive 98/79/EC is available for the national authorities in its facility and that this documentation shall be kept for 5 years after the last product has been manufactured;
- that the manufacturing process follows suitable principles of quality assurance;
- that, has implemented and keep up to date, a post-production surveillance system for monitoring the products;
- that the device in question, was introduced into the market provided with CE mark

Direttore Tecnico/ Technical Director
Dot. Silvio Brocco



PRODOTTI CE DI LIBERA VENDITA / FREE SALE CE PRODUCTS

Rev. 31.0 del 08.01.2016

10002	DNA AGAR + BILI DI TOLLUIDINA	10046	SERUM TELLURITE AGAR
10004	CLED ANDRADE AGAR	10047	BISMUTH SULFITE AGAR
10004*	CLED ANDRADE AGAR	10047*	BISMUTH SULFITE AGAR
10005	MAC CONKEY SORBITOL AGAR	10048	E.M.B. LEVINE AGAR
10005*	MAC CONKEY SORBITOL AGAR	10048*	E.M.B. LEVINE AGAR
10006	TRYPHIC SOY AGAR + 0.6% YEAST EXTRACT	10050	CAMPYLOBACTER AGAR (Sheep Blood 5%)
10007	BACTILUS CEREUS AGAR (PEMBA)	10050*	CAMPYLOBACTER AGAR (Sheep Blood 5%)
10007*	BACTILUS CEREUS AGAR (PEMBA)	10051	Legionella SOYE Agar
10011	YEAST GLUCOSE CHLORAMPHENICOL AGAR	10051*	Legionella SOYE Agar
10011*	YEAST GLUCOSE CHLORAMPHENICOL AGAR	10052	YERSINIA SELECTIVE AGAR
10013	DHES9 TEST AGAR	10052*	YERSINIA SELECTIVE AGAR
10013*	DHES9 TEST AGAR	10053	WILKINS CHALLENGER AGAR
10014	Purple Lactose Agar	10053*	WILKINS CHALLENGER AGAR
10014*	Purple Lactose Agar	10054	WURTZ LACTOSE AGAR
10017	CZAPPEK DOX AGAR	10054*	WURTZ LACTOSE AGAR
10018	DRIGALSKY LACTOSE AGAR	10056*	X.L.D. AGAR
10021	BIGGY (NICKERSON) AGAR	10056*	X.L.D. AGAR
10021*	BIGGY (NICKERSON) AGAR	10057	BILE AESCULIN AGAR
10022	BRIGHTANT GREEN AGAR	10057*	BILE AESCULIN AGAR
10022*	BRIGHTANT GREEN AGAR	10058	TRYPHIC SOY AGAR (Inactivated -30 ml-
10023	Chocobio Agar	10058*	TRYPHIC SOY AGAR (Inactivated -30 ml-
10023*	Chocobio Agar	10060	BRAIN HEART INFUSION AGAR
10025	Chocolate Agar	10060*	BRAIN HEART INFUSION AGAR
10024	TRYPHIC SOY AGAR	10084	CHRISTENSEN UREA AGAR
10024*	TRYPHIC SOY AGAR	10085	SCHAEHLER KKV AGAR(Sheep Blood 5%)
10025	COLLUMBIA AGAR (Horse Blood 5%)	10085*	SCHAEHLER KKV AGAR(Sheep Blood 5%)
10025*	COLLUMBIA AGAR (Horse Blood 5%)	10097	SCHAEHLER KVN AGAR (Sheep Blood 5%)
10026	CLED AGAR	10086	X.L.T. 4 AGAR
10026*	CLED AGAR	10086*	X.L.T. 4 AGAR
10029*	BACTILUS CEREUS AGAR (Mossell)	10074S	TRYPHIC SOY AGAR+NEUTRALIZING Inactivated
10029*	BACTILUS CEREUS AGAR (Mossell)	10074S	TRYPHIC SOY AGAR+NEUTRALIZING Inactivated
10028	ISOSENSITEST AGAR	10078	MUELLER HINTON II MOD AGAR
10028*	ISOSENSITEST AGAR	10078*	MUELLER HINTON II MOD AGAR
10029	ISOSENSITEST AGAR	10079	CASITONE AGAR
10029*	ISOSENSITEST AGAR	10079*	CASITONE AGAR
10029*	MAC CONKEY AGAR	10080	HEMOPHYLUS TEST AGAR
10029*	MAC CONKEY AGAR	10080*	HEMOPHYLUS TEST AGAR
10030	MANNITOL SALT AGAR	10082	HELCOBACTER PYLORI AGAR
10030*	MANNITOL SALT AGAR	10082*	HELCOBACTER PYLORI AGAR
10031	MUELLER HINTON II AGAR	10090	M.R.S. Agar
10031*	MUELLER HINTON II AGAR	10090*	M.R.S. Agar
10033	PSEUDOMONAS ICETRIMIDE YAGAR	10090*	M.R.S. Agar
10033*	PSEUDOMONAS ICETRIMIDE YAGAR	10095	BRAIN HEART AGAR FOR HAEMOPHYLUS
10035	SABOURAUD AGAR	10128	MAC CONKEY AGAR M1G
10035*	SABOURAUD AGAR	10128*	MAC CONKEY AGAR M1G
10035S	SABOURAUD AGAR (Irradiated	10129*	MAC CONKEY AGAR M1G
10035S*	SABOURAUD AGAR (Irradiated	10131	Muller-Hinton II Agar (Sheep Blood 5%)
10036	S.S. AGAR	10131*	Muller-Hinton II Agar (Sheep Blood 5%)
10036*	S.S. AGAR	10132	MUELLER HINTON FASTIDIOUS AGAR 90 mm
10037	TRYPHIC SOY AGAR	10134	Legionella BIPA Agar
10037*	TRYPHIC SOY AGAR	10141	SALMONELLA TEST AGAR
10037S	TRYPHIC SOY AGAR (Inactivated	10141*	SALMONELLA TEST AGAR
10039	FOGOSA AGAR	10142	BLOOD AGAR (Sheep Blood 7%)(ISO 10560)
10039*	FOGOSA AGAR	10142*	BLOOD AGAR (Sheep Blood 7%)(ISO 10560)
10040	NEW YORK CITY AGAR	10143	Muller-Hinton Agar + 5 % Horse Blood lysed
10040*	NEW YORK CITY AGAR	10143	Muller-Hinton Agar + 5 % Horse Blood lysed
10041	LISTERIA PALCAM AGAR	10146	CAMPYLOBACTER PRESTON AGAR
10041*	LISTERIA PALCAM AGAR	10146	CAMPYLOBACTER PRESTON AGAR
10042	CRYSTAL VIOLET AGAR (Sheep Blood 5%)	10148	CAMPYLOBACTER AGAR (Sheep Blood 10%)
10042*	CRYSTAL VIOLET AGAR (Sheep Blood 5%)	10225	LISTERIA PALCAM AGAR 140 mm
10043	HEKTOEN ENTERIC AGAR	10231	MUELLER HINTON II AGAR 140 mm
10043*	HEKTOEN ENTERIC AGAR	10233	R.P.M.I. AGAR
10044	NUTRIENT AGAR	10235	SABOURAUD CAF AGAR + GENTAMICIN
10044*	NUTRIENT AGAR	10235*	SABOURAUD CAF AGAR + GENTAMICIN
10044*	NUTRIENT AGAR	10236	CLED AGAR 140 mm

PRODOTTI CE DI LIBERA VENDITA / FREE SALE CE PRODUCTS

Rev. 31.0 del 08.01.2016

10240	SCHAEDLER K AGAR (Sheep Blood 5%) 140mm
10241	SCHAEDLER KKK AGAR (Sheep Blood 5%) 140mm
10242	SABOURAUD CAF AGAR 140 mm
10243	SABOURAUD CAF AGAR + GENTAMIN 140mm
10244	DERMATOPHYTE (D.T.M.) AGAR 140 mm
10245	BRUCELLA BLOOD AGAR w/ HEMIN AND VITAMIN K1
10246	Chromatic™ MH
10248	Bacterial Blood Agar with Hemin and Vitamin K1
10334	Neomycin Blood Agar (Sheep Blood 5%)
10334*	NEOMYCIN BLOOD AGAR (Sheep Blood 5%)
10335	MUELLER HINTON CHOCOLATE AGAR
10335	BORDET GENGOU AGAR (Sheep Blood 15%)
10335*	BORDET GENGOU AGAR (Sheep Blood 15%)
10407	SCHAEDLER CNA AGAR (Sheep Blood 5%)
10408	WILKINS CHAGREN AGAR + 5% SHEEP BLOOD
10409	CAMPYLOBACTER CODA AGAR
10410	MUELLER HINTON AGAR w/ VITALEX
10411	BLE ESQUILINE AZIDE AGAR w/ VANCOMYCIN
10412	Legionella BCYE Agar w/ Cycloheximide
10413	XLD Agar BP USP, JP Formulation
10416	MINDI EMBROCK TH11 AGAR
10424	Legionella BCYE Agar w/ Vancomycin + Colistin
10425	SCHEOSPORIUM SELECTIVE AGAR
10438	Mycology Agar No. 2
10439	Mycology Agar No. 2
10439*	Group A Selective Strip Agar w/ 5% Sheep Blood
10499	CHROMATIC™ MRS-A
10500	OXA-LINIC RESISTANCE STAPHYLOCOCCUS AGAR
10501	CHOCOLATE AGAR w/ YITOX
10502	HEMOPHAGOCYTES SKIRROW AGAR
10505	HEMOPHAGOCYTES PYCLO EGG YOLK EMULSION
10505	O. LISTERIA
11023	CHOCOLATE BACTRACIN AGAR
11023*	CHOCOLATE BACTRACIN AGAR
11024	COLUMBIA CNA AGAR (Sheep Blood 5%)
11024*	COLUMBIA CNA AGAR (Sheep Blood 5%)
11025	COLUMBIA CNA AGAR (Sheep Blood 5%)
11025*	COLUMBIA AGAR (Sheep Blood 5%)
11027	DESOSYCHOLATE AGAR
11027*	DESOSYCHOLATE AGAR
11030	ANAEROBIC AGAR
11033	PSEUDOMONAS ISOLATION AGAR
11033*	PSEUDOMONAS ISOLATION AGAR
11035	SABOURAUD CAF AGAR
11035*	SABOURAUD CAF AGAR
11038S	SABOURAUD CAF AGAR (Inactivated)
11037	TRYPIC SOY AGAR (Sheep Blood 5%)
11037*	TRYPIC SOY AGAR (Sheep Blood 5%)
11038	TRYPIC SOY AGAR (Horse Blood 5%)
11038*	TRYPIC SOY AGAR (Horse Blood 5%)
11040	THAYER MARTIN AGAR
11040*	THAYER MARTIN AGAR
11041	AZIDE AGAR (Sheep Blood 5%)
11041*	AZIDE AGAR (Sheep Blood 5%)
11052	DERMATOPHYTE (D.T.M.) AGAR
11052*	DERMATOPHYTE (D.T.M.) AGAR
11054	GARDNERELLA AGAR (Sheep Blood 5%)
11054*	GARDNERELLA AGAR (Sheep Blood 5%)

11057	ENTEROCOCCUS AGAR
11057*	ENTEROCOCCUS AGAR
11058	SLANETZ BARTLEY AGAR (ENTEROCOCCUS)
11058*	SLANETZ BARTLEY AGAR (ENTEROCOCCUS)
11060	CLOSTRIDIUM AGAR (Sheep Blood 5%)
11060*	CLOSTRIDIUM AGAR (Sheep Blood 5%)
11067	CLOSTRIDIUM AGAR (Sheep Blood 5%)
11065*	SCHAEDLER K AGAR (Sheep Blood 5%)
11065*	SCHAEDLER K AGAR (Sheep Blood 5%)
11070	HYGOSCEL AGAR
11070*	MYCOSCEL AGAR
11124	COLUMBIA CNA MOD AGAR (Sheep Blood 5%)
11124*	COLUMBIA CNA MOD AGAR (Sheep Blood 5%)
11124*	SABOURAUD AGAR MODIFIED
11135*	SABOURAUD AGAR MODIFIED
11143	HERSEL LEA AGAR
11143*	HERSEL LEA AGAR
11145*	VOGEL JOHNSON AGAR
11145*	VOGEL JOHNSON AGAR
11195*	T.C.B.S. AGAR
11195*	T.C.B.S. AGAR
11196	SPS AGAR
11196*	SPS AGAR
11200*	PAR TEST AGAR
11200*	PAR TEST AGAR
11205	MYCOPLASMA AGAR
11206	Mueller Hinton II Agar (Sheep Blood 5%) 140mm
11206*	Mueller Hinton II Agar (Sheep Blood 5%) 140mm
11235	SABOURAUD CAF AGAR + TTC
11235*	SABOURAUD CAF AGAR + TTC
11236	Sabouraud CAF Agar + Additive
11250	TINSDALE AGAR
11250*	TINSDALE AGAR
11335*	SABOURAUD AGAR + GENTAMIN
11335*	SABOURAUD AGAR + GENTAMIN
11506	BURKHOLDERIA CEPHALIA SELECTIVE AGAR
11509	R.P.M.I. AGAR
11510	M.HINTON+GLUCOSE-METHYLEN BLUE
11511	NERSEERIA/MORAXELLA MEDIUM
11512	NUTRIENT AGAR acc to ISO 21528
11513	NUTRIENT AGAR acc to ISO 6579
11517	COLUMBIA AGAR/Sheep Blood 5%+ VANCOMYCIN
11518	Mueller Hinton Agar + Cloxacillin
11610	Chromatic™ E coli O157
11611	CHROMATIC™ DETECTION
11612	CHROMATIC™ CANDIDA
11614	CHROMATIC™ S.A. MONELLA
11616	CHROMATIC™ STAPH AUREUS
11617	CHROMATIC™ STREPTO B
11618	CHROMATIC™ M.H.
11619	CHROMATIC™ GRE
11621	CHROMATIC™ Y/E
11622	CHROMATIC™ ESSEL
11627	Chromatic™ Enterococcus
11629	CHROMATIC™ ESSEL + AmpC
11629*	CHROMATIC™ ESSEL + AmpC
11631	Chromatic™ OXA-48
11632	Chromatic™ Clostridium difficile
11634	Chromatic™ Detraction plaque

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12031	MUELLER HINTON II AGAR (120x120 mm)
12032	Mueller Hinton II Agar (Sheep Blood 5%) (120 mm x 120 mm)
12033	Mueller Hinton Fertilisation Agar (Horse Blood 5% + 20 mg/ml β-NAD) (120 mm x 120 mm)
13012	CLEDMACCONKEY VISA BLOOD AGAR
13012*	CLEDMACCONKEY VISA BLOOD AGAR
13012*	CLEDMACCONKEY VISA BLOOD AGAR
13013	BAIRD PARKER/SPS/MACCONKEY
13013*	BAIRD PARKER/SPS/MACCONKEY
13014	COLUMBIA DUBOCCO/CLONTHAYER MARTIN
13014*	COLUMBIA DUBOCCO/CLONTHAYER MARTIN
13014*	COLUMBIA DUBOCCO/CLONTHAYER MARTIN
13017*	CLEDMACCONKEY V MANGALTO
13018	BROM CRESOL PURPLE/COLUMBIA CNA/M CONKEY
13018*	BROM CRESOL PURPLE/COLUMBIA CNA/M CONKEY
13018*	BROM CRESOL PURPLE/COLUMBIA CNA/M CONKEY
13018*	CLEDMACCONKEY/CFE/IRIMIDE
13018*	CLEDMACCONKEY/CFE/IRIMIDE
13019*	MAC CONKEY/B PARKER/ISA BLOOD
13020	MAC CONKEY/B PARKER/ISA BLOOD
13045	GARDNERELLA V. ROGOSSA/THAYER MARTIN
13045*	GARDNERELLA V. ROGOSSA/THAYER MARTIN
13056	Group V / Cholesterol / Thayer Martin
13071*	BAIRD PARKER/MACCONKEY/SABOURAUD CAF
13071*	BAIRD PARKER/MACCONKEY/SABOURAUD CAF
13071*	MAC CONKEY/VOGEL JOHNSON/SABOURAUD
13071*	MAC CONKEY/VOGEL JOHNSON/SABOURAUD
13082	SABOURAUD C/CF/BAIRD PARKER/BILE ESQUILINE
13082*	SABOURAUD C/CF/BAIRD PARKER/BILE ESQUILINE
13087	CHOC. BAC./COLUMBIA/MAC CONKEY
13087*	CHOC. BAC./COLUMBIA/MAC CONKEY
13087*	CLEDMACCONKEY/ENTEROCOCCO
13087*	CLEDMACCONKEY/ENTEROCOCCO
13087*	MYCOPLASMA AGAR
18007	CHROMATIC™ STAPH AUREUS / MRS-A
18008	TSB BLOOD/CROMAGAR ORIENTATION
18008*	TSB BLOOD/CROMAGAR ORIENTATION
18008	Chromatic™ Salmonella/Hidran Enteric
18012	CHROMATIC™ DETECTION/ESSEL
18012*	CHROMATIC™ DETECTION/ESSEL
18012*	BRILLIANT GREEN /SS AGAR
18015	BIGGY NICKERSON / MALT AGAR
18015*	BIGGY NICKERSON / MALT AGAR
18017	COLUMBIA CNA BLOOD/CHROMAGAR
18017*	COLUMBIA CNA BLOOD/CHROMAGAR
18018	MAC CONKEY/SABOURAUD CAF
18020	EMB LEVINE / TSA BLOOD
18020*	EMB LEVINE / TSA BLOOD
18021	Chromatic™ GRE/ Chromatic™ ESSEL
18021*	Chromatic™ GRE/ Chromatic™ ESSEL
18022	TSA Blood/Columbia CNA
18027	COLUMBIA CNA/MAC CONKEY
18027*	COLUMBIA CNA/MAC CONKEY
18037	GARDNERELLA V. / THAYER MARTIN
18037*	GARDNERELLA V. / THAYER MARTIN
18380	MAC CONKEY / TSA BLOOD
18380*	MAC CONKEY / TSA BLOOD
18390	BAIRD PARKER / SABOURAUD CAF
18390*	BAIRD PARKER / SABOURAUD CAF
18391	HEKTOEN ENTERIC / YERSINIA
18391*	HEKTOEN ENTERIC / YERSINIA
18422	COLUMBIA CNA / GARDNERELLA
18422*	COLUMBIA CNA / GARDNERELLA

19500	BAIRD PARKER / MAC CONKEY
19500*	BAIRD PARKER / MAC CONKEY
19502	CLEB / MAC CONKEY
19502*	CLEB / MAC CONKEY
19503	HEKTOEN ENTERIC / SS
19503*	HEKTOEN ENTERIC / SS
19505	MAC CONKEY / S. AGAR
19505*	MAC CONKEY / S. AGAR
19507	COLUMBIA CNA / CHOCOLATE
19507*	COLUMBIA CNA / CHOCOLATE
19507*	D.T.M. / SABOURAUD
19507*	D.T.M. / SABOURAUD
19700	Group A Selective TSA II - Sheep Blood 5%
18703	CHOCOLATE AGAR THAYER MARTIN
20075	MAC CONKEY BROT (7/18/20/21/20/6/1)
20077	PHYSIOLOGICAL SOLUTION 2.5 ml
20079	PHYSIOLOGICAL SOLUTION 4.5 ml
20081	INOCULUM SOLUTION 5 ml
20089	SUSPENSION BROTH
20090	HELICOBACTER PYLORI TEST
20095	PHYSIOLOGICAL SOLUTION
20098	PEPTONE WATER
20105	Guinea Bish
20101	INOCULUM BROTH 7 ML
20109	TRYPIC SOY BROTH 15 ml
20109*	TRYPIC SOY BROTH 15 ml
20109*	TRYPIC SOY BROTH 15 ml
20109*	PURE LACTOSE BROTH
20109*	SUSPENSION MEDIUM 7 ML
20158	MYCOPLASMA TRANSPORT BROTH
20159	TRICHOMONAS BROTH w/ CLOSTRIDIUM/HEMICOL
20171	Thioglycolate Medium w/ Vit K1 & Hemin
20340	VAGNIELE
21104	TRYPIC SOY BROTH
21110	SEL ENITE BROTH
21241	Fluid Thioglycolate Medium
21300	SCHAEDLER BROTH
23001	F.B. FASTIDIOUS BROTH
23002	MUELLER HINTON BROTH w/ HORSE BLOOD (11ml)
24070	MUELLER HINTON BROTH
24071	Coxsack Meat Medium
24091	HAEMOPHILUS TEST BROTH 20 PV
24098	PEPTONE WATER 20PV
24100	ALKALINE PEPTONE WATER 20PV
24103	NUTRIENT BROTH 20PV
24104	BRAIN HEART INFUSION BROTH 20PV
24105	Giucose Broth
24107	MUELLER HINTON II BROTH 20 PV
24109	MULLER KAUFMANN BROTH 20PV
24110	SABOURAUD BROTH (Hem E) 20PV
24111	SEL ENITE BROTH 20PV
24111	TOOD HEWITT BROTH 20PV
24112	TRYPIC SOY BROTH 20PV
24115	TRICHOMONAS BROTH 20PV
24117	Fragsile Broth
24119	GM HAJNA BROTH 20PV
24120	BLE ASCULIN BROTH 20PV
24124	Fluid Thioglycolate Medium
24125	SERUM BROTH 20PV
24127	Fluid Thioglycolate Medium + 1% Tween 80

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24128	TRYPTIC SOY BROTH + TWEEN 80 1%	20PV
24136	SALMONELLA DIFFERENTIAL BROTH	20PV
24139	TRYPANE WATER	20PV
24137	MAALONATE BROTH	20PV
24138	LYSINE DECARBOXYLASE BROTH	20PV
24141	BRAIN HEART INFUSION BROTH 2ml	20PV
24142	PHYSIOLOGICAL SOLUTION 3ml	20PV
24144	TODD HEWITT w. Gentamidine acid	20PV
24145	TODD HEWITT B. w. Gentamidine &	20PV
24146	THIOGLYCOLATE M w/o INDICATOR	see USP 20PV
24147	Thioglycollate Bile	20PV
24148	MR-YIP MEDIUM	20PV
24161	Seroumum Deactose Broth + CAF	
24241	Fluid Thioglycollate Medium	
24342	MOTILITY TEST MEDIUM	20PV
24345	O F Medium with Glucose	
24400	RAPPAPORT VASSILADIS SOY (RSV) BROTH	20PV
24404	CAMPYLOBACTER BROTH	20PV
24411	S.F. BROTH	20PV
24412	STREPTOCOCCUS BROTH	20PV
24413	MOSSEL AND MARTIN w/ MANNITOL	20PV
24416	UREA BROTH	20PV
24417	Wilms Chaperon Broth	20PV
24430	SCHAEDLER BROTH	20PV
24432	VERNSIA BROTH	20PV
24433	EUGON BROTH	20PV
24436	WINDLEBROOK THB BROTH	20PV
24446	PHENOL RED BROTH	20PV
24450	Rappaport Broth w/ Soy	
24451	T deactivante Broth	
24459	CASO BROTH (Double Concentration) CE	20PV
24461	RPM Broth	
24462	RPM Broth (double strength)	
24514	TRYPIC SOY BROTH (Hem EP)	
24516	UREA BROTH	
25105	Glucose Broth	
25124	Fluid Thioglycollate Medium - 100 x 1.0 ml	
25129	RAPPAPORT VASSILADIS SOY (RSV) BROTH	
25133	Triple Soy Broth	
27001	GEISA MEDIUM	
27500	Triple Soy Broth	
27501	Todd Hewitt Broth	
27502	Brain Heart Infusion Broth	
27503	Nutrient Broth	
29000	CHECK SET BROTH (Inactivated - 20 T tests)	
30008	CAMPYLOBACTER SELECTIVE THIOGLYCOLATE MEDIUM	
30009	CLOSTRIDIUM AGAR (Sheep Blood 5%)	
30010	HELIODACTER P.V. ORI AGAR	
30011	STREPTOCOCCAL MP + TTC AGAR	
30013	SAMONIS CITRATE AGAR	
30014	NITRATI AGAR	
30016	MOSSEL AGAR	
30022	T.C.B.S. AGAR	
30024	SAROUAUD CAF + ACTIDIONE AGAR	
30030	M.R.S. AGAR	
30080	BORDET GENGOU AGAR (Sheep Blood 15%)	
30081	CHRISTENSEN UREA AGAR	

30082	TRYPTIC SOY AGAR	
30083	NUTRIENT AGAR	
30084	BRAIN HEART INFUSION AGAR	
30085	PHENYLALANINE AGAR	
30087	RIGLER RGN AGAR	
30088	RIGLER RGN AGAR + NaCl 2%	
30090	Muller Hinton II Agar	
30091	BIGGY (NICKERSON) AGAR	
30093	SAROUAUD AGAR	
30095	SIM MEDIUM	
30096	T.S.I. AGAR	
30097	Thydesa Agar	
30098	LYSINE RNOV AGAR	
30099	Chocobite Agar	
30116	LOEFLER MEDIUM	
30117	PEROGIA MEDIUM	
30118	Lowenstein Jensen Medium	
30119	LOWENSTEIN JENSEN MEDIUM w/ G. YC-FEROL	
30121	Sheridink Medium	
30125	DORSET EGG MEDIUM	
30388	MODLEBROOK TH10 AGAR	
31095	SFS Agar	
31075	Muller Hinton II Agar	
31090	Muller Hinton II Agar	
31092	Thydesa Agar	
31099	Chocobite Agar	
31121	Sheridink Medium	
30040	THAYER MARTIN AGAR	
30985	MOSSEL AGAR	
30060	SERUM TELLURITE AGAR	
30086	O.N.P. & AGAR	
30085	BILE AESCULIN AGAR	
30086	DERMATHOPHYTE (D.T.M.) AGAR	
33118	L.I.T.M. MEDIUM	
33120	PETRAKIAN MEDIUM	
34071	CYSTINE TRYPTIC AGAR (CTA)	
34075	Muller Hinton II Agar	
34121	LOWENSTEIN JENSEN + RIFAMPICIN 15 µg/ml	
34121/1	LOWENSTEIN JENSEN + RIFAMPICIN 5 µg/ml	
34121/2	LOWENSTEIN JENSEN + RIFAMPICIN 10 µg/ml	
34121/3	LOWENSTEIN JENSEN + RIFAMPICIN 25 µg/ml	
34121/4	LOWENSTEIN JENSEN + RIFAMPICIN 50 µg/ml	
34121/5	LOWENSTEIN JENSEN + RIFAMPICIN 40 µg/ml	
34121/6	LOWENSTEIN JENSEN + RIFAMPICIN 20 µg/ml	
34122	LOWENSTEIN JENSEN + RIFAMPICIN 9 µg/ml	
34123	LOWENSTEIN JENSEN + ISONIAZID 0.1 µg/ml	
34123/1	LOWENSTEIN JENSEN + ISONIAZID 0.2 µg/ml	
34123/2	LOWENSTEIN JENSEN + ISONIAZID 1 µg/ml	
34123/3	LOWENSTEIN JENSEN + ISONIAZID 5 µg/ml	
34123/4	LOWENSTEIN JENSEN + ISONIAZID 10 µg/ml	
34124/1	LOWENSTEIN JENSEN + PIRAZINAMIDE 5 µg/ml	
34124/2	LOWENSTEIN JENSEN + PIRAZINAMIDE 15 µg/ml	
34124/3	LOWENSTEIN JENSEN + PIRAZINAMIDE 20 µg/ml	
34124/4	LOWENSTEIN JENSEN + PIRAZINAMIDE 30 µg/ml	
34124/5	LOWENSTEIN JENSEN + PIRAZINAMIDE 40 µg/ml	
34124/6	LOWENSTEIN JENSEN + PIRAZINAMIDE 50 µg/ml	
34125/5	LOWENSTEIN JENSEN + STRIPTOMYCIN 50 µg/ml	

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34126/1	LOWENSTEIN JENSEN + ETHAMBUTOL 2 µg/ml	
34126/2	LOWENSTEIN JENSEN + ETHAMBUTOL 4 µg/ml	
34126/3	LOWENSTEIN JENSEN + ETHAMBUTOL 5 µg/ml	
34126/4	LOWENSTEIN JENSEN + ETHAMBUTOL 1 µg/ml	
34126/5	LOWENSTEIN JENSEN + ETHAMBUTOL 3 µg/ml	
34126/6	LOWENSTEIN JENSEN + ETHAMBUTOL 10 µg/ml	
34127	LOWENSTEIN JENSEN + AMIKACIN 5 µg/ml	
34127/1	LOWENSTEIN JENSEN + AMIKACIN 40 µg/ml	
34128/1	LOWENSTEIN JENSEN + OFLOXACIN 5 µg/ml	
34128/2	LOWENSTEIN JENSEN + OFLOXACIN 25 µg/ml	
34128/4	LOWENSTEIN JENSEN + PAS 1 µg/ml	
34128/5	LOWENSTEIN JENSEN + PAS 5 µg/ml	
34130/1	LOWENSTEIN JENSEN + RIFABUTIN 10 µg/ml	
34130/2	LOWENSTEIN JENSEN + RIFABUTIN 30 µg/ml	
34130/3	LOWENSTEIN JENSEN + RIFABUTIN 50 µg/ml	
34131/1	LOWENSTEIN JENSEN + CLARITROMICIN 4 µg/ml	
34131/2	LOWENSTEIN JENSEN + CLARITROMICIN 32 µg/ml	
34131/4	LOWENSTEIN JENSEN + ETHIONAMIDE 40 µg/ml	
34132	LOWENSTEIN JENSEN + ETHIONAMIDE 20 µg/ml	
34132/1	LOWENSTEIN JENSEN + NICOTINAMIDE 20 µg/ml	
34132/2	LOWENSTEIN JENSEN + NICOTINAMIDE 30 µg/ml	
34136	LOWENSTEIN JENSEN + BEN OXACIN 2 µg/ml	
34137/1	LOWENSTEIN JENSEN + CYCLOSERINE 30 µg/ml	
34137/2	LOWENSTEIN JENSEN + CYCLOSERINE 10 µg/ml	
34137/3	LOWENSTEIN JENSEN + CYCLOSERINE 20 µg/ml	
34137/4	LOWENSTEIN JENSEN + CYCLOSERINE 40 µg/ml	
34137/5	LOWENSTEIN JENSEN + CYCLOSERINE 50 µg/ml	
34138/1	LOWENSTEIN JENSEN + CAPREOMYCIN 30 µg/ml	
34138/2	LOWENSTEIN JENSEN + CAPREOMYCIN 40 µg/ml	
34138/3	LOWENSTEIN JENSEN + CAPREOMYCIN 20 µg/ml	
34138/4	LOWENSTEIN JENSEN + CAPREOMYCIN 50 µg/ml	
34139/1	LOWENSTEIN JENSEN + CLOFAZIMINE 5 µg/ml	
34139/2	LOWENSTEIN JENSEN + CLOFAZIMINE 10 µg/ml	
34142/1	LOWENSTEIN JENSEN + KANAMYCIN 10 µg/ml	
34142/2	LOWENSTEIN JENSEN + KANAMYCIN 20 µg/ml	
34142/3	LOWENSTEIN JENSEN + KANAMYCIN 30 µg/ml	
34145	LOW. JENSEN + PACT	
34146/1	Lowenstein Jensen + Leoflozacin 2 µg/ml	
35000	LOWENSTEIN JENSEN MEDIUM	
35002	LOWENSTEIN JENSEN + ISONIAZID 0.20 µg/ml	
35010	LOWENSTEIN JENSEN + RIFAMPICIN 40 µg/ml	
35011	LOWENSTEIN JENSEN + RIFAMPICIN 20 µg/ml	
35022	LOWENSTEIN JENSEN + STREPTOMYCIN 4 µg/ml	
35023	LOWENSTEIN JENSEN + STREPTOMYCIN 10 µg/ml	
35030	LOWENSTEIN JENSEN + ETHAMBUTOL 2 µg/ml	
35040	LOWENSTEIN JENSEN + ETHIONAMIDE 30 µg/ml	
35041	LOWENSTEIN JENSEN + ETHIONAMIDE 40 µg/ml	
35050	LOWENSTEIN JENSEN + PIRAZINAMIDE 1 µg/ml	
35060	LOWENSTEIN JENSEN + KANAMYCIN 20 µg/ml	

35061	LOWENSTEIN JENSEN + KANAMYCIN 30 µg/ml	
35070	LOWENSTEIN JENSEN + PAS 1 µg/ml	
35071	LOWENSTEIN JENSEN + PAS 0.5 µg/ml	
35080	LOWENSTEIN JENSEN + OFLOXACIN 2 µg/ml	
35081	LOWENSTEIN JENSEN + OFLOXACIN 10 µg/ml	
35082	LOWENSTEIN JENSEN + OFLOXACIN 40 µg/ml	
35090	LOWENSTEIN JENSEN + CAPREOMYCIN 30 µg/ml	
35091	LOWENSTEIN JENSEN + CAPREOMYCIN 20 µg/ml	
35147	LOWENSTEIN JENSEN + TCH 2 µg/ml	
35148	LOWENSTEIN JENSEN + TCH 2 µg/ml	
36001/1	UTM + STREPTOMYCIN 2 µg/ml	
36001/2	UTM + STREPTOMYCIN 4 µg/ml	
36001/3	UTM + STREPTOMYCIN 10 µg/ml	
36001/4	UTM + STREPTOMYCIN 25 µg/ml	
36001/5	UTM + STREPTOMYCIN 50 µg/ml	
36002/1	UTM + ISONIAZID 0.1 µg/ml	
36002/2	UTM + ISONIAZID 0.2 µg/ml	
36002/3	UTM + ISONIAZID 1 µg/ml	
36002/4	UTM + ISONIAZID 5 µg/ml	
36002/5	UTM + ISONIAZID 10 µg/ml	
36003/1	UTM + ETHAMBUTOL 1 µg/ml	
36003/2	UTM + ETHAMBUTOL 2 µg/ml	
36003/3	UTM + ETHAMBUTOL 3 µg/ml	
36003/4	UTM + ETHAMBUTOL 5 µg/ml	
36003/5	UTM + ETHAMBUTOL 10 µg/ml	
36004/1	UTM + RIFAMPICIN 5 µg/ml	
36004/2	UTM + RIFAMPICIN 10 µg/ml	
36004/3	UTM + RIFAMPICIN 20 µg/ml	
36004/4	UTM + RIFAMPICIN 40 µg/ml	
36004/5	UTM + RIFAMPICIN 50 µg/ml	
36005/1	UTM + RIFABUTIN 10 µg/ml	
36005/2	UTM + RIFABUTIN 20 µg/ml	
36005/3	UTM + RIFABUTIN 30 µg/ml	
36005/4	UTM + RIFABUTIN 40 µg/ml	
36005/5	UTM + RIFABUTIN 50 µg/ml	
36006/1	UTM + CYCLOSERINE 10 µg/ml	
36006/2	UTM + CYCLOSERINE 20 µg/ml	
36006/3	UTM + CYCLOSERINE 30 µg/ml	
36006/4	UTM + CYCLOSERINE 40 µg/ml	
36006/5	UTM + CYCLOSERINE 50 µg/ml	
36007/1	UTM + OFLOXACIN 1.25 µg/ml	
36007/2	UTM + OFLOXACIN 2.5 µg/ml	
36007/3	UTM + OFLOXACIN 5 µg/ml	
36007/4	UTM + OFLOXACIN 25 µg/ml	
36007/5	UTM + OFLOXACIN 50 µg/ml	
36008/1	UTM + PAS 0.1 µg/ml	
36008/2	UTM + PAS 0.5 µg/ml	
36008/3	UTM + PAS 1 µg/ml	
36008/4	UTM + PAS 5 µg/ml	
36008/5	UTM + PAS 10 µg/ml	
36009/1	UTM + PIRAZINAMIDE 10 µg/ml	
36009/2	UTM + PIRAZINAMIDE 30 µg/ml	
36009/3	UTM + PIRAZINAMIDE 50 µg/ml	
36009/4	UTM + PIRAZINAMIDE 90 µg/ml	
37001	MDDI.EBROCK TH11 + AMIKACIN 2 µg/ml	
37002	MDDI.EBROCK TH11 + AMIKACIN 4 µg/ml	
37006	MDDI.EBROCK TH11 + ETHAMBUTOL 7.5 µg/ml	
37011	MDDI.EBROCK TH11 + ETHIONAMIDE 10 µg/ml	

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37016	MIDDLEBROOK 7H11 + BONAVIDE 0.2 µg/ml
37017	MIDDLEBROOK 7H11 + KAMAYTON 6 µg/ml
37021	MIDDLEBROOK 7H11 + TWEEN 1% 6 µg/ml
37026	MIDDLEBROOK 7H11 + PAS 8 µg/ml
37031	MIDDLEBROOK 7H11 + PYRAZINAMIDE 25 µg/ml
37036	MIDDLEBROOK 7H11 + RIFABUTIN 1 µg/ml
37037	MIDDLEBROOK 7H11 + RIFABUTIN 0.5 µg/ml
37041	MIDDLEBROOK 7H11 + RIFAMPICIN 1 µg/ml
37046	MIDDLEBROOK 7H11 + STREPTOMYCIN 2 µg/ml
37051	MIDDLEBROOK 7H11 + OFLOXACIN 30 µg/ml
37056	MIDDLEBROOK 7H11 + CYCLOSERINE 30 µg/ml
400000	Fluid Thioglycollate Medium 6 x 100 ml
400100	Fluid Thioglycollate Medium 6 x 300 ml
400200	Fluid Thioglycollate Medium 6 x 1000 ml
401980	BUFFER SOLUTION pH 7 6X100 ml
401990	SP8 Agar 6X150 ml
401980	TRYPICONE WATER 6X100 ml
401980	ALVALINE PEPTONE WATER 6X100 ml
402000	NUTRIENT BROTH 6X100 ml
402010	MUELLER HINTON II AGAR 6X100 ml
402030	MULLER KAUFMANN BROTH 6X100 ml
402040	SABOURAUD BROTH 6X100 ml
402050	Selenite Broth 6X100 ml
402060	SALMONELLA DIFF BROTH 6X90 ml
402070	TRYPICONE BROTH 6X100 ml
402120	MRS AGAR 6X100 ml
402130	PEPTONE WATER 6X100 ml
402140	BLOOD AGAR BASE 6X100 ml
402170	AZOE BLOOD AGAR BASE 6X100 ml
402180	GLD AGAR 6X100 ml
402190	NUTRIENT AGAR 6X100 ml
402200	DERMATHOPHYTE (D.T.M.) AGAR 6X100 ml
402210	COLUMBIA CNA AGAR BASE 6X100 ml
402220	DRIGALSKI LACTOSE AGAR 6X100 ml
402230	HECTOGEN ENTERIC AGAR 6X100 ml
402240	MAC CONKEY AGAR 6X100 ml
402250	MUELLER HINTON II AGAR 6X100 ml
402270	PSEUDOMONAS CETRIMIDE AGAR 6X100 ml
402280	SABOURAUD AGAR 6X100 ml
402300	S.S. AGAR 6X150 ml
402310	TRYPICONE AGAR 6X100 ml
402330	BRIILLANT GREEN AGAR 6X100 ml
402340	DESOSYCHOLATE AGAR 6X100 ml
402350	E.M.B. LEVINE AGAR 6X100 ml
402360	SALMONELLA RAPID TEST 6X100 ml
402370	SABOURAUD CAF AGAR 6X100 ml
402380	BRAIN HEART INFUSION AGAR 6X100 ml
402390	PERITONE DILUTIONS 6X100 ml
402400	MAC CONKEY SORBITOL AGAR 6X100 ml
402500	Fluid Thioglycollate Medium + 1% Tween 80
402570	X.L.D. AGAR 6X100 ml
403030	BLOTONE BROTH 6X100 ml
403060	UREA INDOLE BROTH 6X100 ml
413010	BRAIN HEART INFUSION BROTH 6X200 ml
413030	SIMMONS CITRATE AGAR 6X200 ml
413040	LYSINE IRON AGAR 6X200 ml
413060	Selenite Broth 6X200 ml
413080	TOOD HEWITT BROTH 6X200 ml

412060	TRICHOMONAS BROTH 6X200 ml
412100	CHRISTENSEN-UREA AGAR 6X200 ml
412110	TRYPIC SOY BROTH + TWEEN 1% 6X200ml
412130	PSEUDOMONAS AGAR BASE 6X200ml
412150	AZIDE BLOOD AGAR BASE 6X200 ml
412170	PHENYLALANINE AGAR 6X200 ml
412190	NUTRIENT AGAR 6X200 ml
412210	COLUMBIA CNA AGAR BASE 6X200 ml
412230	HEKTOGEN ENTERIC AGAR 6X200 ml
412240	MAC CONKEY AGAR 6X200 ml
412250	MUELLER HINTON II AGAR 6X200 ml
412270	PSEUDOMONAS CETRIMIDE AGAR 6X200 ml
412280	SABOURAUD AGAR 6X200 ml
412290	MANNTOL SALT AGAR 6X200 ml
412300	S.S. AGAR 6X200 ml
412370	SABOURAUD CAF AGAR 6X200 ml
413010	ISOSENSITEST AGAR 6X200 ml
413030	CAMPYLOBACTER AGAR 6X200 ml
413040	CLOSTRIDIUM AGAR BASE 6X200 ml
413080	NUTRIENT AGAR acc. to ISO 6578
414010	PEPTONE WATER pH 8.4 + NaCl 1% 6X22.5 ml
432050	SELENITE BROTH (DOUBLE CONCENTR.) 6X200ml
432060	TRYPIC SOY BROTH 6X225 ml
432250	D-NINE TEST AGAR 6X200 ml
432290	TRYPIC SOY AGAR 6X200 ml
442080	TRYPIC SOY BROTH 6X200 ml
442220	Chocobio Agar 6x 100 ml
442260	SABOURAUD MODIFIED AGAR 6X100 ml
442290	TRYPIC SOY AGAR 6X100 ml
442320	WURTZ LACTOSE AGAR 6X100 ml
442350	BILE AESCULIN AGAR 6X100 ml
442360	BIGGY (NICKERSON) AGAR 6X100 ml
442490	SPS AGAR 6X100 ml
462060	Fluid Thioglycollate Medium 6 x 100 ml
462080	TRYPIC SOY BROTH 6X100 ml
462210	COLUMBIA AGAR BASE 6X200 ml
462500	Fluid Thioglycollate Medium + 1% Tween 80 25 x 100 ml
463000	Fluid Thioglycollate Medium 25 x 100 ml
463100	Selenite Broth 6X1000 ml
463130	Selenite Broth 6X1000 ml
470010	TRYPIC SOY AGAR 6X500 ml
470020	Selenite Broth 6X500 ml
470030	DESOSYCHOLATE AGAR 6X500 ml
470040	SABOURAUD AGAR 6X500 ml
470050	NUTRIENT BROTH 6X500 ml
470060	Mueller-Hinton II Agar 6X500 ml
470070	MANNTOL SALT AGAR 6X500 ml
470090	MAC CONKEY AGAR 6X500 ml
470100	COLUMBIA AGAR BASE 6X500 ml
470110	GLD AGAR 6X500 ml
470120	Chocobio Agar 6 x 500 ml
470140	BILE AESCULIN AGAR 6X500 ml
470150	TRICHOMONAS BROTH 6X500 ml
470160	DESOSYCHOLATE CITRATE AGAR 6X500 ml
470210	ALKALINE PEPTONE WATER 6X500 ml
470220	GAZAPR DOX AGAR 6X500 ml
470280	DRIGALSKI LACTOSE AGAR 6X500 ml

PRODOTTI CE DI LIBERA VENDITA / FREE SALE CE PRODUCTS

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470290	CARY BLAIR TRANSPORT MEDIUM 6X500 ml
470300	Fluid Thioglycollate Medium 6 x 500 ml
470320	PEPTONE WATER 6X500 ml
470370	TRYPIC SOY BROTH 6 x 500 ml
471070	SABOURAUD BROTH 6X500 ml
471200	PHYSIOLOGICAL SOLUTION 6X240 ml
473000	PHYSIOLOGICAL SOLUTION 6X500 ml
481110	CHROMATOC™ CANDIDA 6X100 ml
481130	CHROMATOC™ DETECTION 6X100 ml
481140	CHROMATOC™ SALMONELLA 6X100 ml
481150	CHROMATOC™ STAPH ALBENS 6X100 ml
481160	CHROMATOC™ STREP B 6X100ml
482190	Chromocult™ E col 0157 6 x 200 ml
490010	HEMO-AEROBIC culture 6X50 ml
490020	HEMO-AEROBIC culture 6X30 ml
490030	HEMO-AEROBIC culture-β radiata 6X40 ml
490040	HEMO-AEROBIC culture-β radiata 6X60ml
490050	HEMO-AEROBIC culture HEMOTAL 6x9 ml
490060	HEMO-AEROBIC culture MEDIALAT 6x9 ml
495000	Fluid Thioglycollate Medium 6 x 100 ml
496010	TRYPIC SOY BROTH 6X100 ml
496020	Fluid Thioglycollate Medium 6 x 100 ml
500142	URITEST PENITA
500152	URITEST
500182	URITEST M
500702	URITEST EF
500201	VOGITEST
500211	DERMATTEST
500232	URITEST N
500302	URITEST 2
500402	URITEST MALTO
500412	URITEST EC
51014	URITEST PENITA
51015	URITEST
51018	URITEST M
51020	VOGITEST 120 slide
51021	DERMATTEST
51023	URITEST N
51024	URITEST 2
51041	URITEST EC
51044	URITEST EF
51118	URITEST M
51123	URITEST N 500 slide
51130	URITEST 2 500 slide
51140	URITEST MALTO
51170	CE EDMAC CONKEY BILE AESCULIN
52115	CE EDMAC CONKEY/SLANETZ 120 slide
52119	URITEST SF 500 slide
610001	BILE AESCULIN AZIDE AGAR
610002	DEXTROSE AGAR
610005	BLOOD AGAR BASE
610006	BRAIN HEART INFUSION AGAR
610008	BRAIN HEART INFUSION BROTH
610009	BRIILLANT GREEN AGAR
610012	GLD AGAR
6100125	CE D AGAR

610013	COLUMBIA AGAR BASE
6100135	COLUMBIA AGAR BASE
610014	DESOSYCHOLATE AGAR
6100145	DESOSYCHOLATE AGAR
610015	DESOSYCHOLATE CITRATE AGAR
610016	DRIGALSKI LACTOSE AGAR
610019	E.M.B. LEVINE AGAR
610021	HEKTOGEN ENTERIC AGAR
6100215	HEKTOGEN ENTERIC AGAR
610022	G.C. MEDIUM
610023	KILGER IRON AGAR
610024	M.R.S. AGAR (ISO/DIS 15214)
610025	M.R. 9 BROTH (ISO/DIS 15214)
610026	LOWENSTEIN JENSEN MEDIUM
6100265	LOWENSTEIN JENSEN MEDIUM
610027	LYSINE IRON AGAR
610028	MAC CONKEY AGAR
6100285	MAC CONKEY AGAR
610029	MANNTOL SALT AGAR
6100295	MUELLER HINTON AGAR
610030	MUELLER HINTON AGAR
610032	MXP2 BROTH
610033	MUELLER HINTON BROTH
610034	MUELLER KAUFMANN BROTH
610035	MUELLER HINTON BROTH
610036	NUTRIENT AGAR
610037	NUTRIENT BROTH
610038	NUTRIENT BROTH
610039	PEPTONE WATER
610041	PHENYLALANINE AGAR
6100415	PSEUDOMONAS CETRIMIDE AGAR (ISO 6380-1)
610042	SS AGAR (MODIFIED)
6100425	SS AGAR (MODIFIED)
610043	SCHAEFER AGAR BASE
610044	PIRREL LACTOSE AGAR
610046	SIMMONS CITRATE AGAR
610047	MONSIEUR AGAR
610048	AEROMONAS AGAR BASE
610049	LEGIONELLA BOYE AGAR BASE (ISO 11731)
610050	Fluid Thioglycollate Medium
6100505	Fluid Thioglycollate Medium
610051	TOOD HEWITT BROTH
6100515	TOOD HEWITT BROTH
610052	TRYPIC SOY AGAR (HemmEP) 5 KG
6100525	TRYPIC SOY AGAR
610053	TRYPIC SOY BROTH
6100535	TRYPIC SOY BROTH
6100536	TRYPIC SOY BROTH
610056	CLOSTRIDIUM BROTH
6100565	CLOSTRIDIUM BROTH
6100575	MAC CONKEY AGAR No.2 5 KG
610058	X.L.D. AGAR (ISO 6579)
610065	X.L.D. AGAR
610066	BRAIN HEART INFUSION AGAR
6100665	BRAIN HEART INFUSION BROTH
6100666	BRAIN HEART INFUSION BROTH
610070	YEAST GLUCOSE CHLORAMPHENICOL AGAR
6100705	YEAST GLUCOSE CHLORAMPHENICOL AGAR 5 Kg
610071	PSEUDOMONAS AGAR BASE