

Declaration of CE conformity

Avantor Performance Materials B.V. reg. no. 38013066 who is an established manufacturer of Hematology- Reagents, Stains, Controls and Calibrators and products for Histopathology located at:

Teugseweg 20
7418 AM Deventer
The Netherlands

herewith declares the following:

The reagents (see attached list) are labeled with the J.T.Baker® label and have the CE mark on the label where applicable. The devices comply with the In Vitro Diagnostic Medical Devices Directive 98/79/EC and the conformity assessment procedure according to Annex III. The BeneSphera™ 3 Part Diff Analyzer H32 is in compliance with IEC 61010, Safety Requirements for Electrical Equipment for Measurement, Control and Laboratory Use.

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.

Deventer, the Netherlands.

January 6, 2015

Dr. J. Mittendorf
QA & RA Manager

J.T.Baker® product list for CE marked products

Product no.	Product	Pack size
Hematology Analyzer		
2983	BeneSphera™ 3-part Diff Hematology Analyzer H32	1 unit
Clinical Chemistry Analyzer		
2946	BeneSphera™ Clinical Chemistry Analyzer C72	1 unit
Diluents		
3961	Diluid 100 Plus	20 liter
2990.9010PC	Diluid™ 22	10 liter
3954	Diluid 590	20 liter
3969	Diluid 610	20 liter
3430.9020	Diluid Abacus	20 liter
3430.9010	Diluid Abacus	10 liter
3996	Diluid AC 900	20 liter
3476.9020PC	Diluid APR	20 liter
3957	Diluid Azide free	20 liter
2901.9010PC	Diluid BS34	10 liter
3963	Diluid III Diff	20 liter
3963.9010	Diluid III Diff	10 liter
3459.9020	Diluid Erma	20 liter
3419.9020PC	Diluid M5	20 liter
3439.9020PC	Diluid Mindray	20 liter
3483.9020PC	Diluid NR	20 liter
2987.9020PC	Diluid Ruby	20 liter
3832.9020	Diluid/Sheath 3200-4000	20 liter
3976	Diluid ST 1600/2000	20 liter
3495.9010PC	Sheath D	10 liter
3471.9020PC	Sheath Fluid 3000/3500	20 liter
Lyses		
3998	CN-free Lyse Diff AC 900	5 liter
3744	CyMet 1000 CN free	5 liter
2986.0500PE	CyMet™ 22	500 ml
3469.9010PC	CyMet 3000	10 liter
3823.1000	CyMet 3200 CN free	1 liter
3839.5000PC	CyMet 3500	5 liter
3825	CyMet 3500 CN free	5 liter
3773.5000PC	CyMet 4500 CN free	5 liter
3975	CyMet 530+ CN free	10 liter
3971	CyMet 590 CN free	5 liter
3970	CyMet 610 CN free	10 liter
3977	CyMet 610 CN free	5 liter
3445.1000PE	CyMet Abacus Baso	1 liter
3431.1000	CyMet Abacus CN free	1 liter
3444.1000PE	CyMet Abacus EO	1 liter
3479.1000PE	CyMet APR Baso II	1 liter
3417.0500PE	CyMet APR CN free	500 ml
3478.1000PE	CyMet APR EO	1 liter
2950.2500PE	CyMet ASA	2.5 liter
2951.0500PE	CyMet ASB	500 ml
2952.9010PC	CyMet AS CN Free	10 liter
3755	CyMet Automated	5 liter
2982.0500PE	CyMet BS3 CN free	500 ml
2902.1000PE	CyMet BS34 CN Free	1 liter
3968.0500	CyMet III Diff	500 ml
3968	CyMet III Diff	1 liter
3964	CyMet III Diff	5 liter
3511.1000	CyMet III Diff CN free	1 liter
3511.5000	CyMet III Diff CN free	5 liter

3416.0500	CyMet Erma	500 ml
3841.1000PE	CyMet H12 CN Free	1 liter
3853.1000	CyMet H20	1 liter
3425.0500	CyMet KX CN Free	500 ml
2985.1000PE	CyMet LH 53	1 liter
3489.1000PE	CyMet MBA	1 liter
3418.1000PE	CyMet MD(I)	1 liter
2984.1000PE	CyMet MD(I) 53	1 liter
3488.0500PE	CyMet MD(II)	500 ml
3497.0500PE	CyMet MH CN Free	500 ml
3852.1000	CyMet Micro	1 liter
3863.1000	CyMet Micro CN free	1L micros
3441.0500PE	CyMet Mindray	500 ml
3440.0500PE	CyMet Mindray CN Free	500 ml
3484.1000PE	CyMet NR III	1 liter
3486.1000PE	CyMet NR III CN Free	1 liter
3485.1000PE	CyMet NR V	1 liter
2988.5000PC	CyMet Ruby CN Free	5 liter
3480.5000PC	CyMet SF Baso	5L
3481.5000PC	CyMet SF Diff 1	5L
3482.0500PE	CyMet SF Diff 2	500 ml
3775.1000	CyMet ST 1600/2000	1 liter
3759.5000	CyMet ST 1600/2000 CN free	5 liter
3759.1000	CyMet ST 1600/2000 CN free	1 liter
3788	CyMet STX/STL	1 liter
3475.5000PC	LeucoLyse	5 liter
2989.5000PC	LeucoLyse Ruby	5 liter
3077	LyzerGlobin™	500 ml
3769	LyzerGlobin	6 x 15 ml
3771	LyzerGlobin PCE	6 x 15 ml
3513.1000PE	RBCLyse™	1 liter
3518G.1000PE	RBCLyse G	1 liter
3514.0500PE	WBCStabilise™	500 ml
Reticulocyte Reagents		
3493.1000PE	RetiClear™ MHG	1 liter
3774	RetiCount™	30 ml
2953.0210PE	RetiCount AS	210 ml
3777	RetiCount CD	15 x 3.5 ml
3494.0200PE	RetiCount G	200 ml
Cleaners		
3507.9020	Blanking Solution Hgb	20 liter
3947	Blanking Solution 1600/2000	20 liter
3763	DetectoTerge™	5 liter
3766	DetectoTerge	1 liter
2970.0900PE	DetectoTerge BS	900 ml
3917	HypoChlorite	5 liter
3900	ProClean™	5 liter
3768.1000	ProClean	1L micros
3432.1000PE	ProClean Abacus	1 liter
3432.5000	ProClean Abacus	5 liter
3902.0100PE	ProClean CD	100 ml
3862.9020PC	ProClean Extra	20 liter
3862.5000	ProClean Extra	5 liter
3862.1000	ProClean Extra	1 liter
3867.1000PF	ProClean Extra	1L micros
3498.1000PE	ProClean MX5	1 liter
3901	ProClean Plus	100 ml
3442.5000PE	Rinse Mindray	5 liter

Product no.	Product	Pack size
Reagent Packs		
2910	Reagent Pack BS34	1 pack
Hematology Controls and Calibrators		
3427/3428/3429	8-Parameter Control L/N/H	2.5 ml
3463/3464/3465	8-Parameter Control L/N/H	2.5 ml
3701/3702/3703	8-Parameter Control L/N/H	4.5 ml
3746	8-Parameter Control L+N+H	3 x 2.5 ml
3747	8-Parameter Control 4xN	4 x 2.5 ml
3751	8-Parameter Control 1xL+4xN+1xH	6 x 2.5 ml
3633/3634/3635	8-Parameter Control ext L/N/H	2.5 ml
3433/3434/3435	3-Diff Control L/N/H	2.5 ml
3502/3503/3504	3-Diff Control L/N/H	4.5 ml
3466	3-Diff Control 4xL	4 x 2.5 ml
3467	3-Diff Control 4xN	4 x 2.5 ml
3468	3-Diff Control 4xH	4 x 2.5 ml
3421/3422/3423	3-Diff Control ext L/N/H	2.5 ml
3681/3682/3683	5D Control L/N/H	5.0 ml
3684/3685/3686	ADV-Diff Control L/N/H	3.5 ml
3613/3614/3615	BC-Diff 5 Control L/N/H	4.5 ml
3940	Cal Set 1	2 x 2.5 ml
3452/3453/3454	CD-Diff Control L/N/H	3.0 ml
3838	CD-Diff Control 2xL+2xN+2xH	6 x 3.0 ml
3455/3456/3457	K-Diff Control L/N/H	2.5 ml
3424	Platelet Control Ext. value	5 x 3 ml
3693/3694/3695	SF-Diff Control L/N/H	4.5 ml
3698/3699	WBC reduced RBC Control L/H	3.0 ml
3731/3732/3733	XE-Diff Control L/N/H	4.5 ml
3652/3653/3654	XE-RET Control L/N/H	3.0 ml

Product no.	Product	Pack size
Stains and Dyes		
3800.1000PE	Eosin-Y Alcoholic	1 liter
3800.2500PE	Eosin-Y Alcoholic	2.5 liter
3800.9200	Eosin-Y Alcoholic	200 liter
3446.1000PE	Eosin Y 0.5% Aqueous	1 liter
3446.9200	Eosin Y 0.5% Aqueous	200 liter
3856.0100	Giemsa	0.1 liter
3856.0500	Giemsa	0.5 liter
3856.1000	Giemsa	1 liter
3856.2500	Giemsa	2.5 liter
3856.9180ST	Giemsa	180 liter
3870.1000	Hematoxyline (Mayer)	1 liter
3870.2500	Hematoxyline (Mayer)	2.5 liter
3873.1000	Hematoxyline (Harris, Gill II)	1 liter
3873.2500	Hematoxyline (Harris, Gill II)	2.5 liter
3873.9200	Hematoxyline (Harris, Gill II)	200 liter
3879.1000	Leishman	1 liter
3855.0500	May Grünwald	500 ml
3855.1000	May Grünwald	1 liter
3855.2500	May Grünwald	2.5 liter
3554.1000PE	Papanicolaou Solution 2A	1 liter
3554.2500PE	Papanicolaou Solution 2A	2.5 liter
3554.9200PE	Papanicolaou Solution 2A	200 liter
3555.1000PE	Papanicolaou Solution 2B	1 liter
3555.2500PE	Papanicolaou Solution 2B	2.5 liter
3556.1000PE	Papanicolaou Solution 3B	1 liter
3556.2500PE	Papanicolaou Solution 3B	2.5 liter
3556.9200PE	Papanicolaou Solution 3B	200 liter
3876.1000	Shorr	1 liter
3878.1000	Wright	1 liter
Clearing agent		
3905.2500PE	UltraClear™	2.5 liter
3905.5000PE	UltraClear	5 liter
3905.9010PE	UltraClear	10 liter
3905.9200	UltraClear	200 liter
Mounting media		
3921.0500	UltraKitt™	500 ml
3921.0600	UltraKitt	6 x 100 ml
3921.9025ST	UltraKit	25 liter
3882.0500	Mounting Medium High	500 ml
3883.0500	Mounting Medium Low	500 ml
Fixatives		
3933.1000	10% v/v Buffered Formaldehyde	1 liter
3933.5000PC	10% v/v Buffered Formaldehyde	5 liter
3933.9010PE	10% v/v Buffered Formaldehyde	10 liter
3933.9020	10% v/v Buffered Formaldehyde	20 liter
3933.9200	10% v/v Buffered Formaldehyde	200 liter
3880.1000	Bouin's Fixative	1 liter
3869.1200	Cervix Fixative	12 x 125 ml
3884.9010PC	Cytology Fixative LBCM	10 liter
3409.9010	Immuno PBS 20x concentrated	10 liter
3059	PBS, diluting fluid for blood grouping	20 liter
3059.9010PC	PBS, diluting fluid for blood grouping	10 liter

Declaration of Conformity

helena
Biosciences Europe

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	Clauss Fibrinogen 100	55997
5376H	Clauss Fibrinogen 100	55997

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

Full Name: M.J. Stephenson

Title: Managing Director

Signed:



Date: 05 Aug 2013

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Helena Biosciences Europe
Queensway South, Team Valley Trading Estate,
Gateshead, Tyne and Wear, NE11 0SD,
United Kingdom

Declaration of Conformity

Product Manufactured By: Helena Biosciences Europe
Colima Avenue, Sunderland
Tyne & Wear, SR5 3XB, England

Authorized Representative For: Helena Laboratories, Inc
1530 Lindbergh Drive
Beaumont, TX 77704-0752
USA

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.

Group EDMA Classification Code: 1302500100

Product Code and Description: 5185, SARP (Speciality Assayed Reference Plasma, 10 x 1ml)

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

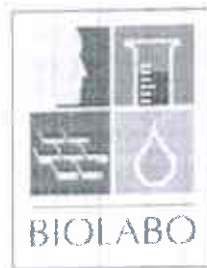
Full Name: M.J. Stephenson

Title: Managing Director

Signed:



Date: 31/10/2003



To whom it may concern

DECLARATION OF EUROPEAN CONFORMITY

LABORATORY REAGENTS & INSTRUMENTS

I, the undersigned, Mrs Oget Isabelle, Regulatory Affairs Director of BIOLABO S.A.S, certify that our Reagents and Instruments (**ACCORDING TO ATTACHED LIST, 5 PAGES**) are manufactured by BIOLABO S.A.S in its Maizy facilities (Les Hautes Rives, F-02160, France) for a world-wide distribution including European Union (EU).

- 1) These products fulfil the essential requirements (Annexe 1) of European Directive IVDM 98/79/CE.
- 2) Essential requirements are reviewed by checking the technical files, including the following information :
 - File for checking Essential Requirements of above mentioned European Directive.
 - File for device's design
 - File for performance (technical specifications).
 - Process management (BIOLABO Standard Operating Procedures, ISO 9001:2008 & 13485:2004 certified)
 - Labelling instructions and references.
 - Package inserts instructions and references.
 - File for batches Traceability including customer's information.
 - Risk Analysis, based on EN ISO 14971.
- 3) BIOLABO S.A.S Quality System Management is ISO 9001:2008 certified under No 1999/12367.10 and ISO 13485:2004 certified under n° 2008/31601.4, by AFAQ (French Association for Quality Assurance).
- 4) I declare that the above information is true and sincere, certifying the product mentioned above fully comply with European Directive 98/79/CE.
- 5) I commit myself to provide to competent French Republic authorities any information which would be requested related to this product, whatever is the origin of such request which may come from their foreign homologues.

This Declaration is issued at Maizy, France, on 20 October 2014.



I. OGET
REGULATORY AFFAIRS DIRECTOR



BIOLABO - Désignation des Dispositifs / Devices Designation

REF	DESIGNATION FR	DESIGNATION GB
80351	ACIDE URIQUE Méthode Uricase	URIC ACID Uricase Method
80001	ACIDE URIQUE Méthode Uricase	URIC ACID Uricase Method
87601	ACIDE URIQUE Méthode Uricase	URIC ACID Uricase Method
80002	ALBUMINE Méthode BGC	ALBUMIN BGC Method
99029	ALCOOL Ethanol	ALCOHOL Ethanol
99059	ALCOOL Ethanol	ALCOHOL Ethanol
80027	ALT/TGP (IFCC.) Monoréactif	ALT/GPT (IFCC.) Single Vial
80127	ALT/TGP (IFCC.) Monoréactif	ALT/GPT (IFCC.) Single Vial
80227	ALT/TGP (IFCC.) Monoréactif	ALT/GPT (IFCC.) Single Vial
80327	ALT/TGP (IFCC.) Monoréactif	ALT/GPT (IFCC.) Single Vial
92027	ALT/TGP Méthode Colorimétrique	ALT/GPT Colorimetric Method
99523	AMYLASE CNPG3	AMYLASE CNPG3
99123	AMYLASE CNPG3	AMYLASE CNPG3
99223	AMYLASE CNPG3	AMYLASE CNPG3
80023	AMYLASE Méthode E-PNPG7	AMYLASE E-PNPG7 Method
80123	AMYLASE Méthode E-PNPG7	AMYLASE E-PNPG7 Method
80223	AMYLASE Méthode E-PNPG7	AMYLASE E-PNPG7 Method
99261	AMMONIAC Méthode Enzymatique	AMMONIA Enzymatic Method
80025	AST/TGO (IFCC.) Monoréactif	AST/GOT (IFCC.) Single Vial
80125	AST/TGO (IFCC.) Monoréactif	AST/GOT (IFCC.) Single Vial
80225	AST/TGO (IFCC.) Monoréactif	AST/GOT (IFCC.) Single Vial
80325	AST/TGO (IFCC.) Monoréactif	AST/GOT (IFCC.) Single Vial
92025	AST/TGO Méthode Colorimétrique	AST/GOT Colorimetric Method
92026	Solution Soude 0,4 N	NaOH Solution 0.4 N
99832	BICARBONATE Méthode Enzymatique	BICARBONATE Enzymatic Method
99852	BICARBONATE Méthode Enzymatique	BICARBONATE Enzymatic Method
80403	BILIRUBINE TOTALE ET DIRECTE Méthode Acide Sulfanilique	TOTAL & DIRECT BILIRUBIN Sulfanilic Acid Method
80443	BILIRUBINE TOTALE Méthode Acide Sulfanilique	TOTAL BILIRUBIN Sulfanilic Acid Method
80553	BILIRUBINE DIRECTE Méthode Acide Sulfanilique	DIRECT BILIRUBIN Sulfanilic Acid Method
97443	BILIRUBINE TOTALE Méthode DCA	TOTAL BILIRUBIN DCA Method
97553	BILIRUBINE DIRECTE Méthode DCA	DIRECT BILIRUBIN DCA Method
90004	CALCIUM Méthode Arsenazo III	CALCIUM Arsenazo III Method
80004	CALCIUM Méthode CPC	CALCIUM CPC Method
80005	CHLORURES Méthode Colorimétrique	CHLORIDE Colorimetric Method
80106	CHOLESTEROL Méthode CHOD-PAP	CHOLESTEROL CHOD-PAP Method
LP80106	CHOLESTEROL CHOD-PAP Liquide Prêt à l'emploi	CHOLESTEROL CHOD-PAP liquid Ready to use
87656	CHOLESTEROL Méthode CHOD-PAP	CHOLESTEROL CHOD-PAP Method
88656	CHOLESTEROL Non estérifié Méthode CHOD-PAP	Non Esterified CHOLESTEROL CHOD-PAP Method
99656	CHOLESTEROL Non estérifié Méthode CHOD-PAP	Non Esterified CHOLESTEROL CHOD-PAP Method
87356	CHOLESTEROL Méthode CHOD-PAP	CHOLESTEROL CHOD-PAP Method
90206	CHOLESTEROL-HDL Méthode Directe	HDL-CHOLESTEROL Direct Method
90406	CHOLESTEROL-HDL Méthode Directe	HDL-CHOLESTEROL Direct Method
90426	CHOLESTEROL-HDL Méthode Directe	HDL-CHOLESTEROL Direct Method
86536	CHOLESTEROL-HDL (PTA) Précipitant	CHOLESTEROL-HDL (PTA) Precipitant
86516	CHOLESTEROL-HDL (PTA) Précipitant	CHOLESTEROL-HDL (PTA) Precipitant
90416	CHOLESTEROL-LDL Méthode Directe	LDL-CHOLESTEROL Direct Method
90816	CHOLESTEROL-LDL Méthode Directe	LDL-CHOLESTEROL Direct Method
82526	CHOLINESTERASE Butyrylthiocholine	CHOLINESTERASE Butyrylthiocholine
97217	Isoenzyme CK - MB Méthode d'immunoinhibition	CK - MB Isoenzyme Immunoinhibition Method
97317	Isoenzyme CK - MB Méthode d'immunoinhibition	CK - MB Isoenzyme Immunoinhibition Method

BIOLABO - Désignation des Dispositifs / Devices Designation

REF	DESIGNATION FR	DESIGNATION GB
92207	CK - NAC IFCC Monoréactif	CK - NAC IFCC Single Vial
92307	CK - NAC IFCC Monoréactif	CK - NAC IFCC Single Vial
80107	CREATININE Méthode cinétique	CREATININE Kinetic method
80008	FER (SFBC) Bathophénanthroline	IRON (SFBC) Bathophenanthrolin
92108	FER Méthode directe (Férène)	IRON Direct Method (Ferene)
92308	C.T.F. Capacité Totale de Fixation du Fer	T.I.B.C. Total Iron Binding Capacity
97408	C.L.F. Capacité Latente de Fixation du Fer	U.I.B.C Unsaturated Iron Binding Capacity
97089	G6-PDH Méthode cinétique U.V.	G6-PDH U.V. Kinetic Method
97099	G6-PDH lyophilisée (méthode U.V.)	Lyophilised G6-PDH U.V. Kinetic Method
80110	GAMMA GT GPNA soluble	GAMMA GT Soluble GPNA
80210	GAMMA GT GPNA soluble	GAMMA GT Soluble GPNA
80310	GAMMA GT GPNA soluble	GAMMA GT Soluble GPNA
81110	GAMMA GT GPNA carboxylé	GAMMA GT carboxy GPNA
81210	GAMMA GT GPNA carboxylé	GAMMA GT carboxy GPNA
81310	GAMMA GT GPNA carboxylé	GAMMA GT carboxy GPNA
80009	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
87109	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
87409	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
16GLB	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
LP80209	GLUCOSE GOD-PAP Liquide Prêt à l'emploi	GLUCOSE GOD-PAP Liquid ready for use
LP87809	GLUCOSE GOD-PAP Liquide Prêt à l'emploi	GLUCOSE GOD-PAP Liquid ready for use
3502200	HEMOGLOBINE Méthode Colorimétrique (Cyanméthémoglobine)	HAEMOGLOBIN Colorimetric Method (Cyanmethemoglobin)
82250	HEMOGLOBINE Méthode Colorimétrique (Cyanméthémoglobine; 1/50)	HAEMOGLOBIN Colorimetric Method (Cyanmethemoglobin; 1/50)
92011	L.D.H. (LDH-P) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
92111	L.D.H. (LDH-P) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
92511	L.D.H. (LDH-P) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
99881	Lipase Méthode cinétique	LIPASE Kinetic method
99891	Lipase Méthode cinétique	LIPASE Kinetic method
87212	MAGNESIUM Calmagite	MAGNESIUM Calmagite
98212	MAGNESIUM CALMAGITE Haute Stabilité - Haute Linéarité	MAGNESIUM CALMAGITE High Stability - High Linearity
92214	PHOSPHATASE ALCALINE (DEA)	ALKALINE PHOSPHATASE (DEA)
92314	PHOSPHATASE ALCALINE (DEA)	ALKALINE PHOSPHATASE (DEA)
82560	PHOSPHATASES ACIDES Méthode Cinétique	ACID PHOSPHATASE Kinetic Method
3300060	PHOSPHATASES ACIDES Méthode Point Final (PNPP)	ACID PHOSPHATASE End Point Method (PNPP)
99105	PHOSPHOLIPIDES Méthode colorimétrique enzymatique	PHOSPHOLIPIDES Colorimetric enzymatic Method
99110	PHOSPHOLIPIDES Méthode colorimétrique enzymatique	PHOSPHOLIPIDES Colorimetric enzymatic Method
80015	PHOSPHORE Inorganique Méthode U.V.	Inorganic PHOSPHOROUS U.V. Method
80016	PROTEINES TOTALES Méthode Biuret	TOTAL PROTEIN Biuret Method
LP87016	PROTEINES TOTALES Méthode Biuret Prêt à l'emploi	TOTAL PROTEIN Biuret Method Ready to use
97016	PROTEINES U.S. Méthode Rouge de Pyrogallol	U.S. PROTEIN Pyrogallol Red Method
80019	TRIGLYCERIDES Méthode GPO	TRIGLYCERIDES GPO Method
87319	TRIGLYCERIDES Méthode GPO	TRIGLYCERIDES GPO Method
80221	UREE Méthode colorimétrique	UREA Colorimetric Method
80321	UREE Méthode colorimétrique	UREA Colorimetric Method
92032	UREE U.V. Méthode Cinétique	UREA U.V. Kinetic Method
92132	UREE U.V. Méthode Cinétique	UREA U.V. Kinetic Method
99032	UREE U.V. Méthode Cinétique Haute Linéarité	UREA U.V. High Linearity Kinetic Method
99132	UREE U.V. Méthode Cinétique Haute Linéarité	UREA U.V. High Linearity Kinetic Method
92315	KIT CALCULS URINAIRES Méthode qualitative chimique	STONE ANALYSIS SET Chemical qualitative method
92330	KIT CALCULS URINAIRES Méthode qualitative chimique	STONE ANALYSIS SET Chemical qualitative method

BIOLABO - Désignation des Dispositifs / Devices Designation

REF	DESIGNATION FR	DESIGNATION GB
95010	BIOLABO EXATROL-N Taux normaux	BIOLABO EXATROL-N Normal values
95011	BIOLABO EXATROL-P Taux pathologiques	BIOLABO EXATROL-P Pathological values
95015	BIOLABO MULTICALIBRATOR Calibrateur Multiparamétrique	BIOLABO MULTICALIBRATOR Multiparametric calibrator
95020	BIOLABO EEQ Evaluation externe de la qualité	BIOLABO EQA External Quality Assessment
95403	BIOLABO CONTROLE PEDIATRIQUE	BIOLABO PAEDIATRIC CONTROL
95406	CALIBRATEUR CHOLESTEROL-HDL	HDL-CHOLESTEROL CALIBRATOR
95806	CALIBRATEUR CHOLESTEROL-LDL	LDL-CHOLESTEROL CALIBRATOR
95506	CALIBRATEUR HDL/LDL/CK-MB	HDL/LDL/CK-MB CALIBRATOR
95516	Sérum de contrôle HDL/LDL/CK-MB/Lipides "Taux 1"	Control serum HDL/LDL/CK-MB/Lipides "Level 1"
95526	Sérum de contrôle HDL/LDL/CK-MB/Lipides "Taux 2"	Control serum HDL/LDL/CK-MB/Lipides "Level 2"
95801	Calibrant LIPASE	LIPASE Calibrator
95013	Contrôle Normal AMMONIAC / ALCOOL / BICARBONATE	Normal Control AMMONIA / ALCOHOL / BICARBONATE
95023	Contrôle Patho. AMMONIAC / ALCOOL / BICARBONATE	Pathological Control AMMONIA / ALCOHOL / BICARBONATE
95089	G6-PDH Contrôle Normal (hémolysat humain lyophilisé)	G6-PDH Normal Control (Lyophilised human hemolysed blood)
95289	G6-PDH Contrôle Déficient (hémolysat humain lyophilisé)	G6-PDH Deficient Control (Lyophilised human hemolysed blood)
95012	Contrôle urinaire "Taux 1" et "Taux 2"	Urinary Control "Level 1" and "Level 2"
95315	KIT CALCULS URINAIRES Contrôles Positifs et Négatifs	STONE ANALYSIS SET Positive and Negative Controls
13880	BIO-TP Temps de Quick Taux de Prothrombine (TP)	BIO-TP Prothrombin Time (PT)
13885	BIO-TP Temps de Quick Taux de Prothrombine (TP)	BIO-TP Prothrombin Time (PT)
13881	BIO-TP Temps de Quick Taux de Prothrombine (TP)	BIO-TP Prothrombin Time (PT)
13702	BIO-TP LI (Low ISI) Temps de Quick Taux de Prothrombine (TP)	BIO-TP LI (Low ISI) Prothrombin Time (PT)
13704	BIO-TP LI (Low ISI) Temps de Quick Taux de Prothrombine (TP)	BIO-TP LI (Low ISI) Prothrombin Time (PT)
13712	BIO-TP LI (Low ISI) Temps de Quick Taux de Prothrombine (TP)	BIO-TP LI (Low ISI) Prothrombin Time (PT)
13883	TAMPON OWREN KOLLER	OWREN-KOLLER BUFFER
13560	BIO-CK Temps de Thromboplastine Partielle activée (TCA)	BIO-CK Activated Partial Thromboplastin Time (APTT)
13570	BIO-CK Temps de Thromboplastine Partielle activée (TCA)	BIO-CK Activated Partial Thromboplastin Time (APTT)
13660	BIO-SIL Temps de Thromboplastine activée (TCA) Méthode silice	BIO-SIL Activated Partial Thromboplastin Time (APTT) Silice Method
13670	BIO-SIL Temps de Thromboplastine activée (TCA) Méthode silice	BIO-SIL Activated Partial Thromboplastin Time (APTT) Silice Method
13565	CHLORURE DE CALCIUM 0.025M	CALCIUM CHLORIDE 0.025M
13450	BIO-FIBRI Dosage Chronométrique du Fibrinogène	BIO-FIBRI Chronometric determination of Fibrinogen
13451	BIO-FIBRI Dosage Chronométrique du Fibrinogène	BIO-FIBRI Chronometric determination of Fibrinogen
13456	TAMPON FIBRINOGENE	FIBRINOGEN BUFFER
13980	BIO-TT Temps de Thrombine	BIO-TT Thrombin Time
13965	TP-CAL/SET SET de Plasmas de Référence Détermination du Taux de Prothrombine (Temps de Quick)	TP-CAL/SET Standard Set For use during the determination of Prothrombin Time (if to be expressed as %)
13970	BIO-CAL Plasma de référence Pour la calibration lors de la détermination du fibrinogène et du Temps de Quick (Taux de Prothrombine %)	BIO-CAL Reference Plasma For use as calibration plasma during the determination of Fibrinogen and Prothrombin Time (if to be expressed as %)
13961	PLASMA CONTRÔLE TAUX NORMAUX	CONTROL PLASMA NORMAL VALUES
13962	PLASMA CONTRÔLE TAUX PATHOLOGIQUES HAUTS	CONTROL PLASMA PATHOLOGICAL HIGH VALUES
13963	PLASMA CONTRÔLE TAUX PATHOLOGIQUES BAS	CONTROL PLASMA PATHOLOGICAL LOW VALUES
13210	D-Dimer	D-Dimer
13211	D-Dimer Control 1	D-Dimer Control 1
13212	D-Dimer Control 2	D-Dimer Control 2
13302	Factor II - Plasma Déficient	Factor II - Deficient plasma
13305	Factor V - Plasma Déficient	Factor V - Deficient plasma
13307	Factor VII - Plasma Déficient	Factor VII - Deficient plasma

Par le Centre de Recherche et de Développement
 de la Société Française de Chimie Analytique
 et de Biochimie (SFCAB)

BIOLABO - Désignation des Dispositifs / Devices Designation

REF	DESIGNATION FR	DESIGNATION GB
13308	Factor VIII - Plasma Déficient	Factor VIII - Deficient plasma
13309	Factor IX - Plasma Déficient	Factor IX - Deficient plasma
13310	Factor X - Plasma Déficient	Factor X - Deficient plasma
13971	Control N - Plasma de contrôle Niveau 1 pour l'hémostase	Control N - Control plasma level 1 for coagulation
13972	Control P - Plasma de contrôle Niveau 2 pour l'hémostase	Control P - Control plasma level 2 for coagulation
9905TH	S. Typhi H (d.H)	S. Typhi H (d.H)
9905TO	S. Typhi O (9,12-O)	S. Typhi O (9,12-O)
9905AH	S. Paratyphi AH (a-H)	S. Paratyphi AH (a-H)
9905AO	S. Paratyphi AO (1,2,12-O)	S. Paratyphi AO (1,2,12-O)
9905BH	S. Paratyphi BH (b-H)	S. Paratyphi BH (b-H)
9905BO	S. Paratyphi BO (1,4,5-O)	S. Paratyphi BO (1,4,5-O)
9905CH	S. Paratyphi CH (c-H)	S. Paratyphi CH (c-H)
9905CO	S. Paratyphi CO (6,7-O)	S. Paratyphi CO (6,7-O)
9905BA	Brucella abortus	Brucella Abortus
9905PK	Proteus OXK	Proteus OXK
9905P19	Proteus OX19	Proteus OX19
9905P2	Proteus OX2	Proteus OX2
9905BM	Brucella Melitensis	Brucella Melitensis
9905RB	Rose Bengal (B. Abortus)	Rose Bengal (B. Abortus)
9901PC	Contrôle Positif Polyvalent	Positive Polyvalent Control
9901NC	Contrôle Négatif Polyvalent	Negative Polyvalent Control
99054	Antigènes Fébriles Pour Tests de Widal Félix (4x5ml)	Stained Febrile Antigens For Widal Felix Tests (4x5ml)
99056	Antigènes Fébriles Pour Tests de Widal Félix (6x5ml)	Stained Febrile Antigens For Widal Felix Tests (6x5ml)
99058	Antigènes Fébriles Pour Tests de Widal Félix (8x5ml)	Stained Febrile Antigens For Widal Felix Tests (8x5ml)
081050	ASLO-LATEX	ASLO-LATEX
097100	CRP-LATEX	CRP-LATEX
098100	FR-LATEX	FR-LATEX
3800100	RPR-CHARBON	RPR-CHARBON
3800150	RPR-CHARBON	RPR-CHARBON
4500100	TPHA	TPHA
4500200	TPHA	TPHA
085100	HCG-LATEX	HCG-LATEX
RF050E	Facteurs Rhumatoides (RF) - Test Immunoturbidimétrique	RHEUMATOID FACTOR (FR) - Turbidimetric Immunoassay
RF520E	Facteurs Rhumatoides (RF) - Test Immunoturbidimétrique	RHEUMATOID FACTOR (FR) - Turbidimetric Immunoassay
RF/CAL_SET51	BIOLABO RF « Kit de Calibration »	BIOLABO RF « Standard Set »
RF/CAL_SH1	BIOLABO RF Calibrant Super Haut	BIOLABO RF Standard Super High
RF/CONT1	BIOLABO RF Contrôle	BIOLABO RF Control
RF/CONT5	BIOLABO RF Contrôle	BIOLABO RF Control
CRP050E	CRP - Test Immunoturbidimétrique	CRP Turbidimetric Immunoassay
CRP620E	CRP - Test Immunoturbidimétrique	CRP Turbidimetric Immunoassay
CRP/CAL_SET51	BIOLABO CRP « Kit de Calibration »	BIOLABO CRP « Standard Set »
CRP/CAL_SH1	BIOLABO CRP Calibrant Super Haut	BIOLABO CRP Standard Super High
CRP/CONT_L1	BIOLABO CRP Contrôle Bas	BIOLABO CRP Control Low
CRP/CONT_L5	BIOLABO CRP Contrôle Bas	BIOLABO CRP Control Low
CRP/CONT_H1	BIOLABO CRP Contrôle Haut	BIOLABO CRP Control High
CRP/CONT_H5	BIOLABO CRP Contrôle Haut	BIOLABO CRP Control High
ASLO050E	ASLO - Test Immunoturbidimétrique	ASLO Turbidimetric Immunoassay
ASLO620E	ASLO - Test Immunoturbidimétrique	ASLO Turbidimetric Immunoassay
ASLO/CAL_H1	BIOLABO ASLO Calibrant Haut	BIOLABO ASLO Standard High
ASLO/CAL_SH1	BIOLABO ASLO Calibrant Super Haut	BIOLABO ASLO Standard Super High

BIOLABO - Désignation des Dispositifs / Devices Designation

REF	DESIGNATION FR	DESIGNATION GB
ASLO/CAL_SET41	BIOLABO ASLO « Kit de Calibration »	BIOLABO ASLO « Standard Set »
ASLO/CONT1	BIOLABO ASLO Contrôle	BIOLABO ASLO Control
ASLO/CONT5	BIOLABO ASLO Contrôle	BIOLABO ASLO Control
APOA1620E	APOLIPOPROTEINE A1 - Test Immunoturbidimétrique	APOLIPOPROTEINE A1 Turbidimetric Immunoassay
APOB620E	APOLIPOPROTEINE B - Test Immunoturbidimétrique	APOLIPOPROTEINE B Turbidimetric Immunoassay
APOA1050E	APOLIPOPROTEINE A1 - Test Immunoturbidimétrique	APOLIPOPROTEINE A1 Turbidimetric Immunoassay
APOB050E	APOLIPOPROTEINE B - Test Immunoturbidimétrique	APOLIPOPROTEINE B Turbidimetric Immunoassay
A1B/CAL_SET51	BIOLABO A1B « Kit de Calibration »	BIOLABO A1B « Standard Set »
A1B/CAL_H1	BIOLABO A1B Calibrant Haut	BIOLABO A1B Standard High
A1B/CONT1	BIOLABO A1B Contrôle	BIOLABO A1B Control
A1B/CONT5	BIOLABO A1B Contrôle	BIOLABO A1B Control
23010	Microalbumine-Test Immunoturbidimétrique	Microalbumin-Turbidimetric Immunoassay
23011	Microalbumine-Test Immunoturbidimétrique	Microalbumin-Turbidimetric Immunoassay
23012	Microalbumine Calibrant Super Haut	Microalbumin Standard Super High
23013	Microalbumine "Kit de calibration"	Microalbumin "Standard Set"
23014	Microalbumine Contrôle	Microalbumin Control
22010	HbA1c-Test Immunoturbidimétrique	HbA1c--Turbidimetric Immunoassay
22011	HbA1c-Test Immunoturbidimétrique	HbA1c--Turbidimetric Immunoassay
22012	HbA1c "Kit de calibration"	HbA1c "Standard Set"
22013	HbA1c "Kit de contrôle"	HbA1c "Control Set"
KENZA MAX	PHOTOMETRE KENZA MAX BioChemisTry	KENZA MAX BioChemisTry PHOTOMETER
KENZA 120TX	KENZA 120TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 120TX - AUTOMATIC BIOCHEMISTRY ANALYSER
KENZA 240 TX	KENZA 240TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 240TX - AUTOMATIC BIOCHEMISTRY ANALYSER
KENZA 240 ISE	KENZA 240ISE - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE avec module ISE	KENZA 240ISE - AUTOMATIC BIOCHEMISTRY ANALYSER with ISE Module
720 ISE	Module ISE	ISE Module
BIOSOLEA 2	BIO SOLEA 2 - COAGULOMETRE 2 CANAUX	BIO SOLEA 2 - COAGULOMETER 2 CHANNELS
BIOSOLEA 4	BIO SOLEA 4 - COAGULOMETRE 4 CANAUX	BIO SOLEA 4 - COAGULOMETER 4 CHANNELS
SOLEA 100	SOLEA 100 - ANALYSEUR AUTOMATIQUE D'HEMOSTASE	SOLEA 100 - FULL AUTOMATED COAGULATION ANALYSER

Document
 par la direction
 by the...
 ENCHS

МЕДИКЛОН
127276 Москва Ботаническая ул. 35. Т/Ф (495) 231-2272 (499) 502-1214

ПАСПОРТ-СЕРТИФИКАТ ПРОИЗВОДИТЕЛЯ
на «Набор реагентов для определения групп крови человека систем АВО, Резус и Келл» по ТУ-9398-101-51203590-2009 (ЦОЛИКЛОНЫ Анти-А, Анти-В и Анти-АВ)

Наименование: Цоликлон Анти-В во флаконах по 10 мл с синими крышками
Серия: 282211 ОКП: 93 9816
Годен: 1 ноября 2020 г. Объем серии: 10000 мл.
Единица: 100 мл Паспорт: Т-18-11-92 от 27-11-2018

Код	Наименование показателя	Характеристика нормы	Результаты испытаний
1	Внешний вид		
1.1	Цоликлон анти-А	Прозрачная жидкость красного цвета	Соответствует
1.2	Цоликлон анти-В	Прозрачная жидкость синего цвета	Соответствует
1.3	Цоликлон анти-АВ	Прозрачная бесцветная жидкость	Соответствует
2	Серологические свойства		
2.1	Специфичность	Цоликлон анти-В не должен давать агглютинации с эритроцитами групп А(I) и О(I)	Соответствует
2.2	Гемоглинирующая способность	Цоликлон анти-АВ не должен давать агглютинации с эритроцитами группы О(I) агглютинация не должна появляться эритроцитами А I и В с соответствующими цоликлонами	Соответствует 10 секунда
2.3	тип	Поданное 10 сек. после смешивания	Соответствует 1:32 - 1:64
		Тип Цоликлоно анти-А в реакции агглютинации на плазмосети эритроцитами группы А(I) 1:32 - 1:64	Соответствует 1:64
		Тип Цоликлоно анти-В в реакции агглютинации на плазмосети эритроцитами группы В(II) 1:64	Соответствует 1:32 - 1:64
		Тип Цоликлоно анти-АВ в реакции агглютинации на плазмосети эритроцитами групп А(I) 1:32 - 1:64 и В(II) 1:64	Соответствует 1:32 - 1:64

Цоликлон соответствует требованиям ТУ - 9398-101-51203590-2009
Заводской номер
Идентификационный номер
М.С. Орлов

МЕДИКЛОН
127276 Москва Ботаническая ул. 35. Т/Ф (495) 231-2272 (499) 502-1214

ПАСПОРТ-СЕРТИФИКАТ ПРОИЗВОДИТЕЛЯ
на «Набор реагентов для определения групп крови человека систем АВО, Резус и Келл» по ТУ-9398-101-51203590-2009 (ЦОЛИКЛОНЫ Анти-А, Анти-В и Анти-АВ)

Наименование: Цоликлон Анти-А во флаконах по 10 мл с красными крышками
Серия: 282111 ОКП: 93 9816
Годен: 1 ноября 2020 г. Объем серии: 10000 мл.
Единица: 100 мл Паспорт: Т-18-11-91 от 22-11-2018

Код	Наименование показателя	Характеристика нормы	Результаты испытаний
1	Внешний вид		
1.1	Цоликлон анти-А	Прозрачная жидкость красного цвета	Соответствует
1.2	Цоликлон анти-В	Прозрачная жидкость синего цвета	Соответствует
1.3	Цоликлон анти-АВ	Прозрачная бесцветная жидкость	Соответствует
2	Серологические свойства		
2.1	Специфичность	Цоликлон анти-В не должен давать агглютинации с эритроцитами групп А(I) и О(I)	Соответствует
2.2	Гемоглинирующая способность	Цоликлон анти-АВ не должен давать агглютинации с эритроцитами группы О(I) агглютинация не должна появляться эритроцитами А I и В с соответствующими цоликлонами	Соответствует 10 секунда
2.3	тип	Поданное 10 сек. после смешивания	Соответствует 1:32 - 1:64
		Тип Цоликлоно анти-А в реакции агглютинации на плазмосети эритроцитами группы А(I) 1:32 - 1:64	Соответствует 1:64
		Тип Цоликлоно анти-В в реакции агглютинации на плазмосети эритроцитами группы В(II) 1:64	Соответствует 1:32 - 1:64
		Тип Цоликлоно анти-АВ в реакции агглютинации на плазмосети эритроцитами групп А(I) 1:32 - 1:64 и В(II) 1:64	Соответствует 1:32 - 1:64

Цоликлон соответствует требованиям ТУ - 9398-101-51203590-2009
Заводской номер
Идентификационный номер
М.С. Орлов

ООО "Медиклон"

МЕДИКЛОН
127276 Москва, Ботаническая ул. 35, 11Ф (495) 231-2272 (499) 502 1214

П А С П О Р Т - С Е Р Т И Ф И К А Т П Р О И З В О Д И Т Е Л Я
на «Набор реагентов для определения групп крови человека систем АВ0, Резус и Келл» по ТУ-9398-101-51203590-2009 (ЦОЛИКЛОН Анти-Д Супер)

Наименование: Поликлон Анти-Д Супер во флаконах по 10 мл с зелеными крышками

Серия: 281411 ОКП: 93 9816

Годен: 11 декабря 2020 г. Объем серии: 10000 мл

Единица: 100 мл Паспорт: Т-18-11-90 от 19.11.2018

Количество единиц 50

Наименование показателя	Характеристика нормы по ТУ	Результаты испытаний
1. Внешний вид	Прозрачная слезка окрашенная жидкость	Соответствует
2 Серологические свойства		
2.1 Специфичность	Целиклон Анти-Д Супер не должен агглютинировать D(-) эритроциты	Соответствует
2.2 Гематтютинирующая способность	Четкая реакция агглютинации должна наступать в течение 30 сек. после смешивания реагента с D(+ эритроцитами	Соответствует 30 сек
2.3 Титр	Титр Целиклона Анти-Д Супер в реакции агглютинации на прекофе с D(-) эритроцитами 1:32 Титр Целиклона Анти-Д Супер в реакции прямой агглютинации с D(+) эритроцитами в титровальном поле 1:256	Соответствует 1:32 1:256

Соответствует требованиям ТУ 9398-101-51203590-2009

М.С. Директор

ООО "Медиклон"

МЕДИКЛОН
127276 Москва, Ботаническая ул. 35, 11Ф (495) 231-2272 (499) 502-1214

П А С П О Р Т - С Е Р Т И Ф И К А Т П Р О И З В О Д И Т Е Л Я
на «Набор реагентов для определения групп крови человека систем АВ0, Резус и Келл» по ТУ-9398-101-51203590-2009 (ЦОЛИКЛОН Анти-А, Анти-В и Анти-АВ)

Наименование: Поликлон Анти-АВ

Серия: 081211 ОКП: 93 9816

Годен: 1 ноября 2020 г. Объем серии: 10000 мл

Единица: 100 мл Паспорт: Т-18-11-92 от 27.11.2018

Наименование показателя	Характеристика нормы	Результаты испытаний
1. Внешний вид	Прозрачная жидкость красного цвета	Соответствует
1.1 Цоликлон анти-А	Прозрачная жидкость синего цвета	Соответствует
1.2 Цоликлон анти-В	Прозрачная бесцветная жидкость	Соответствует
1.3 Цоликлон анти-АВ	Цоликлон анти-А не должен давать агглютинации с эритроцитами групп В(III) и O(0)	Соответствует
2. Серологические свойства	Цоликлон анти-В не должен давать агглютинации с эритроцитами групп A(II) и O(0)	Соответствует
2.1 Специфичность	Цоликлон анти-АВ не должен давать агглютинации с эритроцитами групп A(II) и O(0)	Соответствует
2.2 Гематтютинирующая способность	Цоликлон анти-АВ не должен давать агглютинации с эритроцитами групп A(II) и O(0)	Соответствует 10 секунд
2.3 Титр	Цоликлонами должно появляться не позднее 10 сек. после смешивания	Соответствует 1:32 - 1:64
	Титр Цоликлона анти-А в реакции агглютинации на плоскости с эритроцитами группы A(II) 1:32 - 1:64	Соответствует 1:64
	Титр Цоликлона анти-В в реакции агглютинации на плоскости с эритроцитами группы B(III) 1:64	Соответствует 1:32 - 1:64
	Титр Цоликлона анти-АВ в реакции агглютинации на плоскости с эритроцитами групп A(II) 1:32 и B(III) 1:64	Соответствует 1:32 - 1:64

Соответствует требованиям ТУ 9398-101-51203590-2009

М.С. Директор




Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	HBs Ag one Version ULTRA CODES: SAG1ULTRA.CE (192 tests) SAG1ULTRA.CE.96 (96 tests) SAG1ULTRA.CE.480 (480 tests) SAG1ULTRA.CE.960 (960 tests) SAG1ULTRA.CE.DB (192 tests)
CLASSIFICATION	ANNEX II – LIST A
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318• DESIGN CERTIFICATE N° 2008 12 0588 ED RELEASED BY EC NOTIFIED BODY N° 0318• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318
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PLACE & DATE OF FIRST ISSUE	MILANO – DECEMBER 2008
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 12/2013



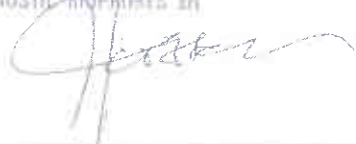
Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

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

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318• DESIGN CERTIFICATE N° 2003 12 0390 ED RELEASED BY EC NOTIFIED BODY N° 0318• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318
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PLACE & DATE OF FIRST ISSUE	MILANO – JANUARY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	 DIA. PRO. DIAGNOSTIC BIOPROBES srl

Rev: 12/2013



Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

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

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318• DESIGN CERTIFICATE N° 2003 12 0391 ED RELEASED BY EC NOTIFIED BODY N° 0318• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318
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PLACE & DATE OF FIRST ISSUE	MILANO – JANUARY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	 DIA. PRO. DIAGNOSTIC BIOPROBES S.R.L.

Rev. 12/2013




Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	HCV Ab CODES: CVAB.CE (192 tests) CVAB.CE.96 (96 tests) CVAB.CE.480 (480 tests) CVAB.CE.960 (960 tests) CVAB.CE.DB (192 tests)
CLASSIFICATION	ANNEX II – LIST A
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318• DESIGN CERTIFICATE N° 2003 12 0392 ED RELEASED BY EC NOTIFIED BODY N° 0318• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318
PLACE & DATE OF FIRST ISSUE	MILANO – JANUARY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	 DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.

Rev: 12/2013



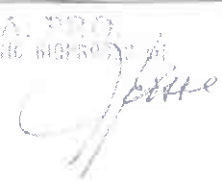
Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

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

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

ISO CERTIFICATE(S)	UNI CEI EN ISO 9001–Nr 50 100 5931/A UNI CEI EN ISO 13485–Nr 50 100 5931/B RELEASED BY CERTIFICATION BODY TÜV Italia S.r.l.
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PLACE & DATE OF FIRST ISSUE	MILANO – MARCH 2004
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	 DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.

Rev: 12/2013



Dia.Pro
Diagnostic
Bio**Probes**

EC DECLARATION OF CONFORMITY

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

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">• FULL QUALITY ASSURANCE SYSTEM N° 2003 12 0388 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318• DESIGN CERTIFICATE N° 2003 12 0393 ED RELEASED BY EC NOTIFIED BODY N° 0318• UNE EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318
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PLACE & DATE OF FIRST ISSUE	MILANO – JANUARY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	 DIA. PRO. DIAGNOSTIC BIOPROBES S.R.L.

Rev. 12/2013

DECLARATION OF CONFORMITY

1) Manufacturer (Name, department): **Monobind Inc.**

Address: **100 North Pointe, LAKE FOREST, CA 92630. UNITED STATES**

and

2) European authorized representative: **CEpartner4U BV,**

Address: **ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS;**

(on product labels printed as:

CEpartner4U , ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS. www.cepartner4u.eu)

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

Immunoassay products;

ELISA,

CLIA,

Control,

Instruments

(see appendix)

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

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

5) Additional information (Conformity procedure, Notified Body, CE certificate, Registration nr., etc.):

Conformity assessment procedure for CE marking: *In vitro* Diagnostic Medical Device Directive, Annex III

Registration nr. : **NL- CA002-22758 and NL- CA002-22762**

Lake Forest, USA; 2013-09-16

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

A Shatola

Tony Shatola; QA Director, Monobind Inc.

(name; function and signature of manufacturer)

Maarn, NL; 2013-09-16

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

Olga Teirlinck

Olga Teirlinck; Consultant, CEpartner4U BV

(name; function and signature of authorized representative)



Appendix

Date: 2013-09-16

List of devices.

Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# QSure® Control	Item# Instrum ent	EDMS code	Risk Class	First date of CE-marking
Thyroid							
Total Triiodothyronine (tT3) Test System	125-300	175-300			12.04.01.05.00	Low	2005-11-11
Free Triiodothyronine (fT3) Test System	1325-300	1375-300			12.04.01.01.00	Low	2005-11-11
Thyroxine (tT4) Test System	225-300	275-300			12.04.01.07.00	Low	2005-11-11
Free Thyroxine (fT4) Test System	1225-300	1275-300			12.04.01.02.00	Low	2005-11-11
Thyrotropin (TSH) Test System	325-300	375-300			12.04.01.11.00	Low	2005-11-11
Rapid TSH Test System	6025-300	6075-300			12.04.01.11.00	Low	2010-06-29
T3-Uptake (T3U) Test System	525-300	575-300			12.04.01.06.00	Low	2005-11-11
Thyroxine-Binding Globulin (TBG) Test System	3525-300	3575-300			12.04.01.09.00	Low	2005-11-11
Thyroglobulin (Tg) Test System	2225-300	2275-300			12.04.01.08.00	Low	2005-11-11
Total Thyroxine (tT4), Total Triiodothyronine (tT3) & Thyroid Stimulating Hormone (TSH) Thyroid Panel (VAST) Test System	8025-300	8075-300			12.04.01.01.00	Low	2005-11-11
Total Triiodothyronine (tT3 SBS) Test System	8125-300	8175-300			12.04.01.01.00	Low	2010-06-29
Total Thyroxine (tT4 SBS) Test System	8225-300	8275-300			12.04.01.01.00	Low	2010-06-29
Free Thyroxine (fT4), Free Triiodothyronine (fT3) & Thyroid Stimulating Hormone Free Thyroid Panel (VAST) Test System	7025-300	7075-300			12.04.01.01.00	Low	2010-06-29
Neonatal Thyroid & Genetics							
Neonatal TSH (N-TSH) Test System	3425-300	3475-300			12.04.01.90.00	Low	2005-11-11
Neonatal (N-T4) Thyroxine Test System	2625-300	2675-300			12.04.01.12.00	Low	2005-11-11
Neonatal 17OHP (N-17OHP) Test System	5525-300	5575-300			12.05.01.07	Low	2008-02-01
Neonatal TBG (N-TBG) Test System	8925-300	8975-300			12.04.01.09.00	Low	2013-09-16
Autoimmune Thyroid							
Anti-Thyroglobulin (Anti-Tg) Test System	1025-300	1075-300			12.10.03.04.00	Low	2005-11-11
Anti-Thyropoxidase (Anti-TPO) Test System	1125-300	1175-300			12.10.03.01.00	Low	2005-11-11
Fertility & Prenatal							
Luteinizing Hormone (LH) Test System	625-300	675-300			12.05.01.05.00	Low	2005-11-11
Folicle Stimulating Hormone (FSH) Test System	425-300	475-300			12.05.01.04.00	Low	2005-11-11
Prolactin Hormone (PRL) Test System	725-300	775-300			12.05.01.08.00	Low	2005-11-11
Prolactin Hormone Sequential (PRLs) Test System	6025-300	6075-300			12.05.01.08.00	Low	2005-11-11
B-Human Chorionic Gonadotropin (hCG) Test System	825-300	875-300			12.05.02.05.00	Low	2005-11-11
B-Human Chorionic Gonadotropin Extended Range (Ext. Range hCG) Test System	8825-300	8875-300			12.05.02.05.00	Low	2013-09-16
Rapid B-Human Chorionic Gonadotropin (Rapid	3325-300				12.05.02.05.00	Low	2005-11-11



Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# QSure® Control	Item# Instrum ent	EDMS code	Risk Class	First date of CE-marking
-hCG) Test System							
Human Chorionic Gonadotropin (hCG) , Human Prolactin (hPRL), Human Luteinizing Hormone (hLH), Follicle Stimulating Hormone (FSH) Fertility Panel (VAST) Test System	8325-300	8375-300			12.05.01.90.00	Low	2006-08-24
Alpha-Fetoprotein (AFP), Human Chorionic Gonadotropin (hCG), Unconjugated Estiol (u-E3) Triple Screen (VAST) Test System	8525-300	8575-300			12.05.01.90.00	Low	2010-06-29
Pregnancy Associated Plasma Protein - A (PAPP-A) Test System	7925-300	7975-300			12.05.02.10.00	Low	2013-09-16
Steroid							
Cortisol Test System	3625-300	3675-300			12.06.02.04.00	Low	2005-11-11
DHEA-S Test System	5125-300	5175-300			12.05.01.02.00	Low	2010-06-29
Dehydroepiandrosterone (DHEA) Test System	7425-300	7475-300			12.05.01.02.00	Low	2011-09-26
Estradiol (E2) Test System	4925-300	4975-300			12.05.01.03.00	Low	2010-06-29
Unconjugated Estiol (u-E3) Test System	5025-300	5075-300			12.05.02.02.00	Low	2010-06-29
Progesterone Test System	4825-300	4875-300			12.05.01.06.00	Low	2010-06-29
Sex Hormone Binding Globulin (SHBG) Test System	9125-300	9175-300			12.05.01.09.00	Low	2013-09-16
Testosterone Test System	3725-300	3775-300			12.05.01.10.00	Low	2007-11-01
Free Testosterone Test System	5325-300	5375-300			12.05.01.10.00	Low	2010-06-29
17α-OH Progesterone Test System	5225-300	5275-300			12.05.01.07.00	Low	2010-06-29
17α-OH Progesterone - SI Test System	9925-300	9975-300			12.05.01.07.00	Low	2010-10-18
Growth & Bone Metabolism							
Growth Hormone (hGH) Test System	1725-300	1775-300			12.06.04.02.00	Low	2005-11-11
Parathyroid Hormone (PTH) Test System	9225-300	9275-300			12.06.03.13.00	Low	2011-09-26
25-Hydroxyvitamin D3 (Vitamin D3) Test System	7725-300	7775-300			12.06.03.10.00	Low	2011-09-26
Diabetes							
Insulin Test System	2425-300	2475-300			12.06.01.03.00	Low	2005-11-11
Rapid Insulin Test System	5825-300				12.06.01.03.00	Low	2010-06-29
C-Peptide Test System	2725-300	2775-300			12.06.01.01.00	Low	2005-11-11
Insulin - C-Peptide (VAST)	7325-300	7375-300			12.06.01.03.00	Low	2005-11-11
Cardiac Markers							
CK-MB Test System	2925-300	2975-300			12.13.01.02.00	Low	2005-11-11
Troponin I (cTnI) Test System	3825-300	3875-300			12.13.01.07.00	Low	2005-11-11
Digoxin (DIG) Test System	925-300	975-300			12.08.01.01.00	Low	2005-11-11
High Sensitivity CRP (hs-CRP) Test System	3125-300	3175-300			12.13.01.90.00	Low	2005-11-11
Myoglobin Test System	3225-300	3275-300			12.13.01.05.00	Low	2005-11-11



Device types	Item# AccuBind® ELISA Microwells	Item# AccuLite® CLIA Microwells	Item# OSure® Control	Item# Instrum ent	EDMS code	Risk Class	First date of CE-marking
Infectious Diseases							
Anti-H. Pylori IgG Test System	1425-300	1475-300			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgM Test System	1525-300	1575-300			15.01.04.03.00	Low	2005-11-11
Anti-H. Pylori IgA Test System	1625-300	1675-300			15.01.04.03.00	Low	2005-11-11
Cancer Markers							
Alpha-Fetoprotein (AFP) Test System	1925-300	1975-300			12.03.90.01.00	Low	2005-11-11
CA-125 Test System	3025-300	3075-300			12.03.01.06.00	Low	2005-11-11
CA 15-3 Test System	5625-300	5675-300			12.03.01.02.00	Low	2010-06-29
CA -19-9 Test System	3925-300	3975-300			12.03.01.03.00	Low	2005-11-11
Carcinoembryonic Antigen (CEA) Test System	1825-300	1875-300			12.03.01.31.00	Low	2005-11-11
Next Generation Carcinoembryonic Antigen (CEA) Test System	4625-300	4675-300			12.03.01.31.00	Low	2010-06-29
Free β-Subunit Human Chorionic Gonadotropin (fβhCG) Test System	2025-300	2075-300			12.03.01.90.00	Low	2005-11-11
Allergy & Anemia							
Ferritin Test System	2825-300	2875-300			12.07.01.02.00	Low	2005-11-11
Folate Test System	7525-300	7575-300			12.07.01.03.00	Low	2010-06-29
Immunoglobulin E (IgE) Test System	2525-300	2575-300			12.02.01.02.00	Low	2005-11-11
Transferrin Soluble Receptor (sTfR) Test System	8625-300	8675-300			12.07.01.06.00	Low	2010-06-29
Vitamin B-12 (B12) Test System	7625-300	7675-300			12.07.02.04.00	Low	2011-09-26
Folate, Vitamin B-12 (VAST) Test System	7825-300	7875-300			12.07.01.00.00	Low	2013-09-16
Miscellaneous Controls							
Anti-Thyroglobulin (Anti-Tg), Anti-Thyroperoxidase (Anti-TPO) Control – Positive & Negative			AIT-101		12.50.01.16.00	Low	2010-06-29
High Level Fertility Control – Single Level – Progesterone, Estradiol, Human Chorionic Gonadotropin			FC-300		12.50.01.16.00	Low	2010-06-29
Maternal Control – Tri Level - Human Chorionic Gonadotropin, Free Beta Human Chorionic Gonadotropin Subunit, Alpha Feta Protein, Estriol			MC-300		12.50.01.16.00	Low	2010-06-29
Thyroglobulin (Tg) Control – Tri Level			TG-300		12.50.01.16.00	Low	2010-06-29
H. Pylori IgG Control – Positive & Negative			HPy-IgG-300		12.50.01.16.00	Low	2010-06-29
H. Pylori IgM Control – Positive & Negative			HPy-IgM-300		12.50.01.16.00	Low	2013-09-16
H. Pylori IgA Control – Positive & Negative			HPy-IgA-300		12.50.01.16.00	Low	2013-09-16
Thyroid Binding Globulin (TBG) Control – Tri-Level			TBG-300		12.50.01.16.00	Low	2013-09-16
Miscellaneous Instruments							
Autoplex ELISA & CLIA Analyzer				IN006	21.02.10.01	Low	2010-06-29
Autoplex Generation 2 ELISA & CLIA Analyzer				IN005-2	21.02.10.01	Low	2013-09-16
Lumax CLIA Analyzer				IN001	21.02.10.01	Low	2008-08-24
Neo-Lumax CLIA Analyzer				IND10	21.02.10.01	Low	2011-09-26



Declaration of Conformity

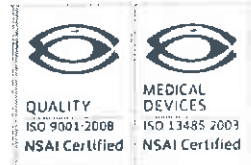
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Page: 5 of 5

<i>Device types</i>	<i>Item# AccuBind® ELISA Microwells</i>	<i>Item# AccuLite® CLIA Microwells</i>	<i>Item# QSure® Control</i>	<i>Item# Instrum ent</i>	<i>EDMS code</i>	<i>Risk Class</i>	<i>First date of CE-marking</i>
Impulse 2 CLIA Analyzer				IN005	21.02.10.01	Low	2006-08-24
Impulse 3 CLIA Analyzer				IN007	21.02.10.01	Low	2010-06-29
Lumax96 CLIA Analyzer				IN004	21.02.10.01	Low	2007-03-01
LuMatic CLIA Analyzer				IN008	21.02.10.01	Low	2011-09-26
Eldex 3.8 ELISA Analyzer				IN003	21.02.10.01	Low	2007-09-10
Neo-Eldex ELISA Analyzer				IN009	21.02.10.01	Low	2011-09-26
PrisMatic ELISA Analyzer				IN013	21.02.10.01	Low	2013-09-16
Plate Washer Microplate Washer				IN002	21.02.10.01	Low	2010-06-29



Monobind, Inc.



DECLARATION OF CONFORMITY

Product Family TOTAL AND FREE PROSTATE SPECIFIC ANTIGEN (PSA and FPSA)

Specific Product Details						
Product Description	Item # ELISA	Item # CLIA	EDMS Code	GMDN ELISA Code	GMDN CLIA Code	Risk Class
Total PSA Immunoassay	2125-300	2175-300	12.03.01.32.00	54664	54665	High/ List B
Free PSA Immunoassay	2325-300	2375-300	12.03.01.33.00	54668	54669	High/ List B
Cancer VAST Immunoassay	8425-300	8475-300	12.03.01.32.00	54664	54665	High/ List B
Multi Ligand Control	ML-300	ML-300	12.03.01.32.00	38207	38207	High/ List B

Manufacturer

Name Monobind Inc.
 Address 100 North Pointe, Lake Forest, CA 92630
 Country United States

Representative

Name CEpartner4U BV,
 Address Esdoornlaan 13, 3951DB Maarn
 Country The Netherlands
 Telephone +31 (0)6 – 516.536.26

Notified Body

Name NSAI
 Body ID Number 0050
 CE Cert # 304.1006
 Registration # NL-CA002-2011-23306

Means of Conformity

Monobind Inc. declares that the product listed is in conformity with the Annex IV, IVD Type List B essential requirements and provisions of Council Directive: 98/79/EC

And is in conformance with the following standards:

EN 13612—2002 EN 980-2008 ISO 14971:2009
 ISO 18113:2009 EN 13641:2002 EN 13640:2002

Under the principles of ISO 13485:2003

Signature

Place and date Monobind Inc. October 28, 2011
 Signature *Ashatola*
 Name Tony Shatola Title QA Director

	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 18113-1:2011; 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

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

No.	Product name	Identification data	REF
1.	Vectohep A-IgG	Enzyme immunoassay kit for the qualitative and quantitative determination of IgG to hepatitis A virus	D-0362
2.	VectoMeasles-IgG	Enzyme immunoassay kit for the quantitative and qualitative determination of IgG to measles virus in blood serum (plasma)	D-1356
3.	VectoMeasles-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.	VectoEBV-NA-IgG	Enzyme immunoassay kit for the detection of IgG to nuclear antigen of Epstein-Barr virus in blood serum (plasma)	D-2170
7.	VectoEBV-EA-IgG	Enzyme immunoassay kit for the detection of IgG to early antigens of Epstein-Barr virus in blood serum (plasma)	D-2172
8.	VectoEBV-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.	VectoMumps-IgG	Enzyme immunoassay kit for the detection of IgG to mumps virus in blood serum (plasma)	D-2602
10.	VectoMumps-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

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

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



SYPHILIS SEROLOGY KIT
DIRECTIONS FOR USE

RPR CARBON KIT: For Detection Of Syphilis.

SUMMARY

At one time, syphilis was a major medical disease with a host of different manifestations transmitted primarily through sexual contact. The advent of penicillin in 1943 changed this. The etiologic agent of syphilis is *Treponema pallidum*, a spiral bacterium (spirochete). The spirochete causes some damage to the heart and the liver, releasing some tissue fragments. The patient's immune system produces antibodies, called reagins, against these fragments. There are two different techniques for the detection of syphilis. TPHA tests, which detect antibodies to *Treponema pallidum*, and non-treponemal serologic tests, which detect Reagin in infected people.

PRINCIPLE

When used by the recommended techniques, the reagent will agglutinate (clump) in the presence of reagin. No agglutination usually indicates the absence of reagin (see **Limitations**).

KIT DESCRIPTION

Lorne RPR Carbon Kit is a non-treponemal serologic test for the detection of syphilis. The RPR Carbon Antigen contains micro particulate carbon, which aids in the microscopic reading of results. All the reagents are supplied at optimum dilution for use with all recommended techniques without the need for further dilution or addition. For lot reference number and expiry date see **Vial Labels**.

STORAGE

Do not freeze. Reagent vials should be stored at 2 - 8°C on receipt. Prolonged storage at temperatures outside this range may result in accelerated loss of reagent reactivity.

SPECIMEN COLLECTION

Specimens should be drawn with or without anticoagulant using an aseptic phlebotomy technique. If testing is delayed specimens can be stored at 2-8°C for 7 days or for up to 3 months at or below -20°C. Specimens must be free from bacterial contamination, fibrin, haemolysis and lipaemia.

PRECAUTIONS

1. The kit is for *in vitro* diagnostic use only.
2. Do not use kit past expiration date (see **Vial and Box Labels**).
3. Protective clothing should be worn when handling the reagents, such as disposable gloves and a laboratory coat.
4. No known tests can guarantee products derived from human or animal sources are free from infectious agents. Care must be taken in the use and disposal of each vial and its contents.
5. RPR Positive Control: H319 - Causes serious eye irritation. Follow the precautionary statement given in the SDS.

DISPOSAL OF KIT REAGENT AND DEALING WITH SPILLAGES

For information on disposal of kit reagent and decontamination of a spillage site see **Material Safety Data Sheets**, available on request.

CONTROLS AND ADVICE

1. It is recommended the RPR Positive and Negative Controls be tested in parallel with each batch of tests. Tests must be considered invalid if controls do not show expected results.
2. Shake all the reagents well before use to ensure homogeneity.
3. Do not interchange components between different kits.
4. The circles on the agglutination cards should never be touched with fingers, as this may invalidate the test results.
5. Use of kit and interpretation of results must be carried out by properly trained and qualified personnel in accordance with the requirements of country where reagents are in use.
6. The user must determine suitability of the kit for use in other techniques.

KIT COMPONENTS PROVIDED

- 1) RPR Carbon Antigen (Red Label): Carbon particles coated with a lipid complex (cardiolipin, lecithin and cholesterol) in phosphate buffer 20 mmol/L, pH 7.0 containing a preservative.
- 2) RPR Positive Control (Red cap): Artificial serum with reagin titer $\geq 1/4$.
- 3) RPR Negative Control (Blue cap): Animal serum containing a preservative
- 4) Dispensing bottle (1 x 2 ml).
- 5) Dispensing Needle (x1).
- 6) Disposable agglutination slides.
- 7) Plastic stirrers.

MATERIALS AND EQUIPMENT NOT SUPPLIED

- a) Pipette capable of accurately delivering 50 μ l
- b) Mechanical rotating table capable of rotating at 80-100 rpm.
- c) 9 g/L saline solution.

QUALITATIVE TECHNIQUE

1. Allow the reagents and samples to reach room temperature. The sensitivity of the test may be reduced at low temperatures.
2. Place 50 μ L of the sample and one drop of each Positive and Negative controls into separate circles on the slide test.
3. Swirl the RPR-carbon reagent gently before using. Invert the dropper assembly and press gently to remove air bubbles from the micropipette.
4. Place the micropipette in a vertical position and perpendicular to the slide, and add one drop (20 μ L) of this reagent next to the samples to be tested.
5. Mix the drops with a stirrer, spreading them over the entire surface of the circle. Use different stirrers for each sample
6. Place the slide on a mechanical rotating table at 80-100 r.p.m. for 8 min. False positive results could appear if the test is read after more than 8 minutes.

INTERPRETATION OF QUALITATIVE RESULTS

1. **Reactive:** Visible agglutination (medium to large clumps) constitutes a positive result and within the accepted limitations of the test procedure, indicates the presence of reagin.
2. **Weak-Reactive:** Weak agglutination (small clumps) around the periphery of the test area constitutes a weak positive result and within the accepted limitations of the test procedure, indicates the presence of reagin.
3. **Negative:** No agglutination constitutes a negative result and within the accepted limitations of the test procedure, indicates the absence of reagin.

SEMI QUANTITATIVE TECHNIQUE

1. The semi-quantitative test can be performed in the same way as the quantitative technique using dilutions of the serum in 9 g/L saline solution.
2. Make doubling dilutions of specimen as follows:

Dilution	Serum	Saline
1/2	100 μ l undiluted serum	100 μ l
1/4	100 μ l 1/2 diluted serum	100 μ l
1/8	100 μ l 1/4 diluted serum	100 μ l
1/16	100 μ l 1/8 diluted serum	100 μ l

3. Test the specimen dilutions in the same way as for the quantitative technique above.
4. Read the test and note the last positive dilution series.

STABILITY OF THE REACTIONS

Slide tests should be interpreted straight after the 8-minute rotating period to avoid the possibility that a negative result may be incorrectly interpreted as positive due to drying of the reagent.

LIMITATIONS

1. RPR carbon test is non-specific for syphilis. All Reactive samples should be retested with treponemic methods such as TPHA and FTA-Abs to confirm the results.
2. A Non Reactive result by itself does not exclude a diagnosis of syphilis. Clinical diagnosis should not be made on findings of a single test result, but should integrate both clinical and laboratory data.
3. False positive results have been reported in diseases such as infectious mononucleosis, viral pneumonia, toxoplasmosis, pregnancy and autoimmune diseases.
4. Bilirubin (≤ 20 mg/dL), hemoglobin (≤ 10 g/L) and lipids (≤ 10 g/L), do not interfere. Rheumatoid factors (≥ 300 IU/mL), interfere. Other substances may interfere⁵.
5. False positive or negative results may also occur due to:
 - a) Not expelling air from end of needle
 - b) Not maintaining dispensing bottle and needle in a vertical position when dispensing the antigen.
 - c) When transferring the specimen from the collecting tube some of the specimen being drawn up in to the test
 - d) Contamination of test materials
 - e) Improper storage of test materials or omission of reagents
 - f) Deviation from the recommended techniques

SPECIFIC PERFORMANCE CHARACTERISTICS

1. The kit has been characterised by all the procedures mentioned in the **Recommended Techniques**.
2. Prior to release, each lot of Lorne RPR Syphilis Kit is tested by the **Recommended Techniques** to ensure suitable reactivity.
3. The reagent sensitivity is calibrated against the "Human Reactive Serum" from the CDC (Centres for Disease Control) and comparable to the RPR reagent from Becton Dickinson.
4. **Prozone effect:** No prozone effect was detected up to titers $\geq 1/128$.
5. **Diagnostic sensitivity:** 100%
6. **Diagnostic specificity:** 100 %.

DISCLAIMER

1. The user is responsible for the performance of the kit by any method other than those mentioned in the **Recommended Techniques**.
2. Any deviations should be validated prior to use using established laboratory procedures.

BIBLIOGRAPHY

1. George P. Schmid. Current Opinion in Infectious Diseases 1994, 7: 34-40.
2. Sandra A Larsen et al. Clinical Microbiology Reviews 1995; 8 (1): 1-21.
3. Sandra Larsen et al. A manual of Test for Syphilis American Public Health Association 1990: 1-192.
4. Joseph Earle Moore et al. Gastrointestinal Haemorrhage 1952; 150(5): 467-473.
5. Young DS. Effects of drugs on clinical laboratory test, 4th ed. AACC Press, 1995.








AVAILABLE KIT SIZES

Kit Size	Catalogue Number
150 Tests Per Kit	044150A
500 Tests Per Kit	044500A

For the availability of other sizes, please contact:

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

TABLE OF SYMBOLS

	Batch Number		In-vitro Diagnostic
	Catalogue Reference		Store At
	Expiry Date		Manufacturer
	Read Pack Insert		

DECLARATION OF CONFORMITY

PRODUCT IDENTIFICATION

Product name	Catalogue number
RPR Carbon kit	044150A 044500A

MANUFACTURER

Name	Lorne Laboratories
Address	Unit 1 Cutbush Park Industrial Estate Danehill Lower Earley Berks, RG6 4UT
Country	United Kingdom

MEANS OF CONFORMITY

I hereby declare that the products listed above comply with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

This declaration is valid from 17 May 2015.



Eddy Velthuis
Technical Director



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

Lorne Laboratories Limited
Unit 1 Cutbush Park Industrial Estate
Danehill, Lower Earley
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CERTIFICATO n. **4265/4/C**
CERTIFICATE No.

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

KIMA S.R.L.

UNITÀ OPERATIVE / OPERATIVE UNITS

Via Leonardo Da Vinci, 22 - Zona Industriale Tognana - 35028 Piove di Sacco (PD)
Italia

È CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

UNI CEI EN ISO 13485:2016

Sistema di Gestione per la Qualità / Quality Management System

PER LE SEGUENTI ATTIVITÀ / FOR THE FOLLOWING ACTIVITIES

EA: 29

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.

*Trading of the products of the Group: diagnostic kits, culture media for microbiology,
plastic disposable labware, test tubes with predetermined vacuum and sterile needles.*

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,
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Data emissione
First issue
18/01/2007

Emissione corrente
Current issue
18/01/2019

Data di scadenza
Expiring date
17/01/2022


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SGQ N° 004 A

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CERTIFICATE No.

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WE HEREBY CERTIFY THAT THE QUALITY MANAGEMENT SYSTEM OPERATED BY

VACUTEST KIMA S.r.l.

Sede / Head office

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

Uffici direzionali e amministrativi

Unità Operative / Operative Units

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

Progettazione e produzione di provette con vuoto predeterminato ad uso 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 predeterminato e aghi sterili.

Via Leonardo Da Vinci, 22 - 35028 Piove di Sacco (PD)

Uffici commerciali e magazzino

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UNI CEI EN ISO 13485:2016

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. 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 predeterminato e aghi sterili.

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. Trading of the products of the Group: diagnostic kits, culture media for microbiology, plastic disposable labware, test tubes with predetermined vacuum and sterile needles.

Riferirsi alla documentazione del Sistema di Gestione per la Qualità aziendale per l'applicabilità dei requisiti della norma di riferimento.
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CISQ is the Italian Federation of management system Certification Bodies.



EG-KONFORMITÄTSERKLÄRUNG • EC DECLARATION OF CONFORMITY

Name und Adresse des Herstellers: Name and address of the manufacturer:	KABE LABORTECHNIK GmbH Jägerhofstraße 17 51588 Nümbrecht-Eisenroth Deutschland / Germany
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Wir erklären in alleiniger Verantwortung, dass die In-Vitro-Diagnostika der Produktgruppe /
We declare under our sole responsibility that the in-vitro-diagnostics of product group


kapillare Blutentnahmesysteme <ul style="list-style-type: none">• Kapillarblutentnahmesystem (GK)• kapillare Probenbehältnisse<ul style="list-style-type: none">• Blutgaskapillaren (BK)• Hämatokritkapillaren (HK)• end-to-end Kapillaren (EK)	capillary blood collection systems <ul style="list-style-type: none">• capillary blood collection system (GK)• capillary sample containers<ul style="list-style-type: none">• blood gas capillaries (BK)• haematocrit capillaries (HK)• end-to-end capillaries (EK)
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der Klasse / of class	Andere IVD-Produkte Other IVD-devices
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den einschlägigen Bestimmungen der IVD-Richtlinie 98/79/EG und deren Umsetzungen in nationale Gesetze entspricht. Die Konformitätserklärung gilt für die durch die KABE LABORTECHNIK GmbH freigegebenen Chargen.
meets the provisions of the directive 98/79/EEC and its transpositions in national laws which apply to it. This declaration is valid for the batches released by KABE LABORTECHNIK GmbH.

Konformitätsbewertungsverfahren: Conformity assessment procedure:	Richtlinie 98/79/EWG Anhang III Directive 98/79/EEC Annex III
--	--

Nümbrecht-Eisenroth, 21.03.2013

KABE LABORTECHNIK GmbH
Jägerhofstraße 17
D-51588 Nümbrecht-Eisenroth
☎ +49 (0) 2203 / 596

André Kolpe, Geschäftsführer / Managing director



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 contents 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

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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° 122D
SGA N° 0200 ISP N° 075E
PRS N° 097C

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