



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	TOXO IgM CODE: TOXOM.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

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

NOTIFIED BODY (EC) CERTIFICATE(S)	AEMPS – n° 0318 <ul style="list-style-type: none">FULL QUALITY ASSURANCE SYSTEM N° 2004 05 0442 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318UNI 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 – MAY 2004
PLACE & DATE OF CURRENT EMISSION	SESTO SAN GIOVANNI (MI) – MAY 2018
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 05/2018



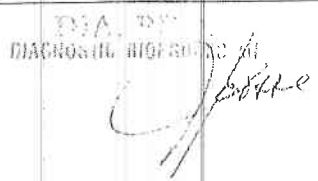
Dia.Pro
Diagnostic
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EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	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	

Rev: 12/2013



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EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	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(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 – APRIL 2004
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev: 12/2013



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EC DECLARATION OF CONFORMITY

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L. VIA G. CARDUCCI N° 27 – 20099 SESTO SAN GIOVANNI (MILANO) – ITALY
PRODUCT	VCA IgM CODE: VCAM.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 – SEPTEMBER 2004
PLACE & DATE OF CURRENT ISSUE	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
SIGNATURE Legal Representative Dr.ssa Fiorenza Scozzesi	

Rev. 12/2013

GBG-MDL SRL
Global Biomarketing Group
Moldova
65 Tighina Str., office 607
MD-2001 Chisinau
Republic of Moldova

NovaTec Immundiagnostica GmbH
Waldstraße 23 A6
63128 Dietzenbach, Germany
Tel.: +49 (0) 60 74 48 76-0
Fax: +49 (0) 60 74 48 76-29
E-Mail: info@NovaTec-ID.com
Internet: www.NovaTec-ID.com

November 18th, 2019

To whomever it may concern:

By hereby, we, **NovaTec Immundiagnostica GmbH**, Waldstraße 23 A6, 63128 Dietzenbach, Germany, declare that **GBG-MDL SRL** Global Biomarketing Group Moldova, 65 Tighina Str., office 607, MD-2001 Chisinau, Republic of Moldova is our authorized distributor designated by us to promote, to register, to distribute and to represent our products in the territory of the Republic of Moldova.

This declaration is valid until December 31st, 2020 unless terminated by either party before the date of expiration.

With best regards

NovaTec Immundiagnostica GmbH


NOVATEC
Britta Maria Duchmann Berlic
General Manager IMMUNDIAGNOSTICA GmbH
Waldstraße 23 A6
63128 Dietzenbach, Germany

Geschäftsführung:
Britta-Maria Duchmann Berlic

Handelsregister: HRB Offentbach 12095

Deutsche Bank
BLZ 50070024
Kto.-Nr. 0106120
BIC: DEUTDE33HAN
IBAN: DE 20 5007 0024 0010 6120 00

Sparkasse Langen-Seligenstadt
BLZ 506 521 24
Kto.-Nr. 5124 300
BIC: HELADEF13LS
IBAN: DE 40 5065 2124 0005 1243 00

Product List – CE Marked

Certified by

ISO 13485:2016

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

2019-10

NovaLisa®

Virology

Prod. No.	Name
ADVA0010	Adenovirus IgA
ADVG0010	Adenovirus IgG
ADVM0010	Adenovirus IgM
CHIG0590	Chikungunya Virus IgG capture
CHIM0590	Chikungunya Virus IgM-µ-capture
CMVG0110	Cytomegalovirus (CMV) IgG
ACMV7110	Avidity Cytomegalovirus (CMV) IgG
CMVM0110	Cytomegalovirus (CMV) IgM
DENG0120	Dengue Virus IgG
DENIM0120	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
HEVG0780	Hepatitis E Virus (HEV) IgG
HEVM0780	Hepatitis E Virus (HEV) IgM
HSVIG0250	Herpes simplex Virus 1+2 (HSV) IgG
HSVIM0250	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
MUMG0340	Mumps Virus IgG
MUMM0340	Mumps Virus IgM
PAIA0360	Parainfluenza Virus 1,2,3 IgA
PAIG0360	Parainfluenza Virus 1,2,3 IgG
PARG0370	Parvovirus B 19 IgG
PARM0370	Parvovirus 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

RUBM0400	Rubella Virus IgM µ-capture
TICG0440	TBE / FSME IgG
TICM0440	TBE / FSME IgM
PTTCG044	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

MYCG0350	Mycoplasma pneumoniae IgG
MYCM0350	Mycoplasma pneumoniae IgM
TETG0430	Clostridium tetani toxin IgG
TETG0443	Clostridium tetani toxin 5S IgG
PTEIG043	Clostridium tetani toxin 5S IgG plus

Parasites

Prod. No.	Name
CHAG0560	Chagas (Trypanosoma cruzi) IgG
TRYP0570	Chagas
ENTG0140	Entamoeba histolytica IgG
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

Bacteriology

Prod. No.	Name
BAR0900	Bartonella
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
PCORR009	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

Prod. No.	Name
HELA0220	Helicobacter pylori IgA
HELG0220	Helicobacter pylori IgG
PHELA022	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

Worms

Prod. No.	Name
ASCG0020	Ascaris lumbricoides IgG
ECHG0130	Echinococcus IgG
FIL0760	Filariasis
SCHG0410	Schistosoma mansoni IgG
SCHM0410	Schistosoma mansoni IgM
STRO0690	Strongyloides
TAEG0420	Taenia solium IgG
TOCG0450	Toxocara canis IgG
TRIG0480	Trichinella spiralis IgG

Fungi

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

NovoLisa® Hormones

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

Prod. No.	Name
ATG1010	Anti-TG
ATPO1020	Anti-TPO
FT41050	Free T4
TSH1030	TSH

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

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

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

Prod. No.	Name
DNOV051	Free T3
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
DNOV 060	CEA
DNOV061	CA 125
DNOV062	CA 15-3
DNOV063	CA 19-9

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

Prod. No.	Name
DNOV100	Ferritin
DNOV101	HGH
DNOV102	IgG

NovoLisa® 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

NovoLisa® Recombinant Antigens

Prod. No.	Name
BORG0040	Borrelia burgdorferi IgG
BORM0040	Borrelia burgdorferi IgM
CHAG0560	Chagas (Trypanosoma cruzi) IgG
TRYP0570	Chagas
HANG0670	Hantavirus IgG
HANM0670	Hantavirus IgM
HELA0220	Helicobacter pylori IgA
PHELA022	Helicobacter pylori IgA plus
HEVG0780	Hepatitis E Virus (HEV) IgG
HEVM0780	Hepatitis E Virus (HEV) 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
MAL0620	Malaria
STRO0690	Strongyloides
ZVG0790	Zika Virus IgG capture
ZVM0790	Zika Virus IgM µ-capture

NovoLisa® 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

NovoLisa® 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
FT41050	Free T4
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

NovoLisa® IgM µ-capture Assays

Prod. No.	Name
CHIM0590	Chikungunya Virus IgM µ-capture
DYM0640	Dengue Virus IgM µ-capture
RUBM0400	Rubella Virus IgM µ-capture
TOXM0460	Toxoplasma gondii IgM µ-capture
ZVM0790	Zika Virus IgM µ-capture

NovoLisa® Antibody Assays

Prod. No.	Name
ASCG0020	Ascanis lumbricoides IgG
CHAG0560	Chagas (Trypanosoma cruzi) IgG
TRYP0570	Chagas
ENTG0140	Entamoeba histolytica IgG
LEIG0310	Leishmania infantum IgG
MAL0620	Malaria
STRO0690	Strongyloides
TAEG0420	Taenia solium IgG
TOCG0450	Toxocara canis IgG
TRIG0480	Trichinella spiralis IgG

NovoLisa® 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

NovoLisa® Liquor Diagnostic

Prod. No.	Name
BORG0040	Borrelia burgdorferi IgG
BORM0040	Borrelia burgdorferi IgM

ВЕКТОР



ОГРН 1025404347550
ИНН 5433104384/ КПП 543301001
р/с 40702810244020101090
в Сибирском банке ПАО Сбербанк,
БИК 045004641
корр. сч. 30101810500000000641
ОКВЭД 21.20.2, 72.11 / ОКПО 23548172

от 18.11.2019 № КВ.151

АО "Вектор-Бест"
630117, г. Новосибирск, а/я 492
тел.: (383) 227-73-60, 332-36-34
тел./факс: 332-67-49, 332-67-52
e-mail: vbmarket@vector-best.ru
Internet: http://www.vector-best.ru

«GBG-MLD» SRL
Республики Молдова, г. Кишинев,
ул. Тигина, 65, оф. 607
Чайковскому Т.К.

Авторизация от производителя

Акционерное общество «Вектор-Бест» (Российская Федерация, 630559, Новосибирская область, рабочий поселок, Кольцово, Научно-производственная зона, корпус 36, к. 211) официально удостоверяет, что «GBG-MLD» SRL (Республика Молдова, г. Кишинев, ул. Тигина, 65, оф. 607) имеет право участвовать от имени АО «Вектор-Бест» в тендерах, проводимых медицинскими учреждениями Республики Молдова и осуществлять рекламно-информационное сопровождение.

Срок действия авторизации по 31 декабря 2020 года включительно.

Коммерческий директор АО «Вектор-Бест»

 Гусев Ю.М.



Сертификат

mdc medical device certification GmbH
удостоверяет, что на предприятии



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

Изделия медицинские – Системы менеджмента качества –
Регулирующие системные требования

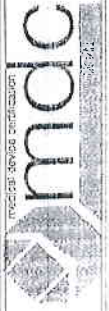
EN ISO 13485:2016 + AC:2016 – ISO 13485:2016

Дата выдачи	2018-07-13
Срок действия до	2020-07-03
Регистрационный №	D1213100017
Отчет №	P18-00489-117996
Штутгарт, Германия	2018-07-13

Руководитель сертификационного органа



mdc medical device certification GmbH
Kriegerstraße 6
D-70191 Stuttgart, Germany
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Fax: +49-(0)711-253597-10



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Fax: +49-(0)711-253597-10

Приложение к Сертификату № D1213100017 от 2018-07-13 Стр. 1 из 1	
Месторасположение АО «Вектор-Бест», ул. Арбузова, 44, 630117, г. Новосибирск, Российская Федерация	Область Действия проектирование и разработка, производство и реализация медицинских изделий in-vitro диагностики
АО «Вектор-Бест», 530559, 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

EC DECLARATION OF CONFORMITY

ZAO "Vector-Best" hereby ensures under own responsibility and declares that the products listed on pages 2-4 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.

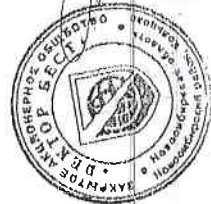
Classification of products: Other devices (all devices except Annex II and self-testing devices)

Conformity assessment procedure: Annex III (not including section 6).

Manufacturer:
ZAO "Vector-Best"
Address: AHC, Koltsovo,
Novosibirsk Region, 630559, Russia,
Tel: +7 (383) 363 20 60,
Fax: +7 (383) 363 35 55

European authorized representative:
Bioron GmbH,
Rheinhorstr. 18, D-67071
Ludwigshafen, Germany.
tel.: +49 (0) 621 5720 915,
fax: +49 (0) 621 5720 916

Date: 2013/04/12



Murat Khusainov
General Director ZAO «Vector-Best»

No.	Product name	Identification data	REF
1.	Vectohep A-IgM	ELISA kit for determination of IgM to hepatitis A virus	D-0352
2.	Vectohep A-IgG	ELISA kit for quantitative and determination of IgG to hepatitis A virus	D-0362
3.	Vectohep TTV-IgG	ELISA kit for determination of IgG to TT virus	D-0302
4.	Vectohep E-IgG	ELISA kit for determination of IgG to hepatitis E virus	D-1056
5.	Vectohep E-IgM	ELISA kit for determination of IgM to hepatitis E virus	D-1058
6.	Vectohep G-IgG	ELISA kit for determination of IgG to hepatitis G virus	D-1252
7.	LymeBest-IgG	ELISA kit for determination of IgG to infectious borreliosis agents	D-1452
8.	LymeBest-IgM	ELISA kit for determination of IgM to infectious borreliosis agents	D-1454
9.	RecombiBest antipallidum-IgG	ELISA kit for determination of IgG to Treponema pallidum	D-1852
10.	RecombiBest antipallidum-total antibodies	ELISA kit for determination of total antibodies to Treponema-pallidum	D-1856
11.	RecombiBest antipallidum-IgM	ELISA kit for determination of IgM to Treponema pallidum	D-1858
12.	RecombiBest antipallidum-total antibodies	ELISA kit for determination of total antibodies to Treponema pallidum	D-1857
13.	VectoHSV-1,2 - IgG	ELISA kit for determination of IgG to herpes simplex virus types 1 and 2	D-2152
14.	VectoHSV - IgM	ELISA kit for determination of IgM to herpes simplex virus types 1 and 2	D-2154
15.	VectoHHV-8 - IgG	ELISA kit for determination of IgG to human herpes virus type 8	D-2160
16.	VectoHHV-6 - IgG	ELISA kit for determination of IgG to human herpes virus type 6	D-2166
17.	Ureaplasma urealyticum - IgG-EIA-BEST	ELISA kit for determination of IgG to Ureaplasma urealyticum antigens	D-2254
18.	Ureaplasma urealyticum - IgA-EIA-BEST	ELISA kit for determination of IgA to Ureaplasma urealyticum antigens	D-2258
19.	VectoParotitis-IgG	ELISA kit for determination of IgG to parotitis virus	D-2602
20.	VectoParotitis-IgM	ELISA kit for determination of IgM to parotitis virus	D-2604
21.	Toxocara-IgG-EIA-BEST	ELISA kit for determination of IgG to toxocara antigens	D-2752
22.	Opisthorchiasis - IgG-EIA-BEST	ELISA kit for determination of IgG to opisthorchiasis antigens	D-2952
23.	Echinococcus-IgG-EIA-BEST	ELISA kit for determination of IgG to Echinococcus	D-3356

24.	Ascarid-IgG-EIA-BEST	antigens ELISA kit for determination of IgG to Ascaris lumbricooides	D-3452
25.	Lambliia-antibodies-EIA-BEST	ELISA kit for determination of IgG, IgM and IgA to Lambliia antibodies	D-3552
26.	Lambliia-IgM-EIA-BEST	ELISA kit for determination of IgM to Lambliia antibodies	D-3554
27.	Lambliia-antigen-EIA-BEST	ELISA kit for determination of Lambliia antigen	D-3556
28.	Helicobacter pylori-CagA-antigen-EIA-BEST	ELISA kit for determination of total antibodies to CagA Helicobacter pylori	D-3752
29.	TSHEIA-BEST	ELISA kit for determination of concentration of thyroid-stimulating hormone	X-3952
30.	T3 total-EIA-BEST	ELISA kit for determination of concentration of total triiodothyronine	X-3954
31.	T4 total-EIA-BEST	ELISA kit for determination of concentration of total thyroxine	X-3956
32.	Anti-TPO-EIA-BEST	ELISA kit for determination of antibody concentration to thyroperoxidase	X-3968
33.	PAPP-A-EIA-BEST	ELISA kit for determination of concentration of pregnancy-associated plasma protein A	D-4160
34.	Mycoplasma hominis-IgG-EIA-BEST	ELISA kit for determination of IgG to Mycoplasma hominis	D-4352
35.	Mycoplasma hominis-IgA-EIA-BEST	ELISA kit for determination of IgA to Mycoplasma hominis	D-4356
36.	Mycoplasma pneumoniae-IgG-EIA-BEST	ELISA kit for determination of IgG to Mycoplasma pneumoniae	D-4362
37.	Mycoplasma pneumoniae-IgM-EIA-BEST	ELISA kit for determination of IgM to Mycoplasma pneumoniae	D-4366
38.	Vectocrimean - CHF - IgG	ELISA kit for determination of IgG to Crimean-Congo hemorrhagic fever virus	D-5052
39.	Vectocrimean - CHF - IgM	ELISA kit for determination of IgM to Crimean-Congo hemorrhagic fever virus	D-5054
40.	CEA-EIA-BEST	ELISA kit for determination of concentration of carcinoembryonic antigen	T-8454
41.	AFP-EIA-BEST	ELISA kit for determination of concentration of Alpha-Fetal Protein	T-8456
42.	CA-125-EIA-BEST	ELISA kit for determination of concentration of oncomarker CA-125	T-8466
43.	CA-19-9-EIA-BEST	ELISA kit for determination of concentration of CA-19-9	T-8470
44.	CA 15-3-EIA-BEST	ELISA kit for determination of concentration of oncomarker CA 15-3	T-8472
45.	NSE-EIA-BEST	ELISA kit for determination of concentration of neuron specific enolase	T-8476

46.	Ferritin-EIA-BEST	ELISA kit for determination of concentration of ferritin	T-8552
47.	IgE total-EIA-BEST	ELISA kit for determination of concentration of total IgE	A-8660
48.	IgG total-EIA-BEST	ELISA kit for determination of concentration of total IgG	A-8662
49.	IgM total-EIA-BEST	ELISA kit for determination of concentration of total IgM	A-8664
50.	IgA total-EIA-BEST	ELISA kit for determination of concentration of total IgA	A-8666
51.	Gamma-Interferon-EIA-BEST	ELISA kit for determination of concentration of gamma-interferon	A-8752
52.	Interleukine-4-EIA-BEST	ELISA kit for determination of concentration of Interleukine-4	A-8754
53.	Alpha-TNF-EIA-BEST	ELISA kit for determination of concentration of alpha-tumor necrosis factor	A-8756
54.	Alpha-Interferon-EIA-BEST	ELISA kit for determination of concentration of alpha-interferon	A-8758
55.	Interleukine-6-EIA-BEST	ELISA kit for determination of concentration of Interleukine-6	A-8768
56.	Interleukine-2-EIA-BEST	ELISA kit for determination of concentration of Interleukine-2	A-8772
57.	Procalcitonin-EIA-BEST	ELISA kit for determination of concentration of procalcitonin	A-9004
58.	NTproBNP-EIA-BEST	ELISA kit for determination of concentration of N-terminal prohormone of brain natriuretic peptide	A-9102
59.	Troponin I-EIA-BEST	ELISA kit for determination of concentration of troponin I	A-9106

Orange County, California, January 10, 2020

IM Global Biomarketing Group - Moldova SRL,
Tighina str.65,office 607
MD-2001,Chisinau, Republic of Moldova

Commercialization Agreement

To Whom It May Concern:

We, Monobind Inc., an ISO 13485 certified company specializing in the research, development and manufacturing of in vitro diagnostic products for clinical and research application, located at 100 North Pointe Drive, Lake Forest, California 92630 USA;

Hereby authorizes and entitles IM Global Biomarketing Group from Moldova legally registered at Tighina str.65,office 607 MD-2001,Chisinau to effect clinical trials and evaluation of goods, registration of the goods at Health Ministry of Moldova, receive certificate of registration and conclude an agreement on consulting and examination of the documents needed for the registration in Moldova.

This is also to confirm that IM Global Biomarketing Group is the exclusive distributor our AccuBind® ELISA and AccuLite® CLIA products and accessories in Moldova. IM Global Biomarketing Group is authorized to promote and supply our products, to contract for their delivery and take part in tenders with our products.

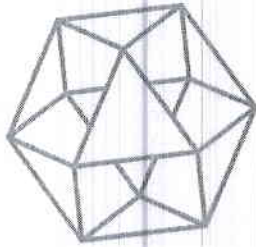
This authorization is valid until January 1, 2021.

On behalf of the Monobind Inc.



Alicia Jerome Volkov
Marketing Director
Monobind Inc.





NSAI

Certificate of Registration of Quality Management System to I.S. EN ISO 13485:2012

The National Standards Authority of Ireland certifies that:

Monobind Inc.
100 North Pointe Drive
Lake Forest, CA 92630
USA

has been assessed and deemed to comply with the requirements of the above standard in respect of the scope of operations given below:

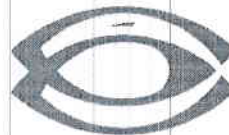
The Design, Manufacture and Distribution of In-Vitro Diagnostic Medical Device Immunoassays, and Related Reagent Products and Equipment

Additional sites covered under this multi-site certification are listed on the Annex (File No. MD19.4585)

Approved by:
Geraldine Larkin
Chief Executive Officer

Approved by:
Susan Murphy
European Medical Device
Operations Manager

Registration Number: MD19.4585
Certification Granted: May 18, 2010
Effective Date: Oct 29, 2017
Expiry Date: Oct 28, 2020



National Standards Authority of Ireland, 1 Swift Square, Northwood, Santry, Dublin 9, Ireland T +353 1 807 3800

DECLARATION OF CONFORMITY

1) **Manufacturer (Name, department):** Monobind Inc.
 Address: 100 North Pointe, LAKE FOREST, CA 92630. UNITED STATES

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:**
 Title: Document No. 98/79/EC
 In vitro Diagnostic Medical Devices Directive

5) **Additional information (Conformity procedure, Notified Body, CE certificate, Registration nr., etc.):**
 Conformity assessment procedure for CE marking: In-vitro Diagnostic Medical Device Directive,
 Annex III
 Registration nr. : NL- CA002-22758 and NL- CA002-22762

Lake Forest, USA; 2013-09-16
 Tony Shatola, QA Director, Monobind Inc.
 (name, function and signature of manufacturer)
 Maarn, NL; 2013-09-16
 Olga Teifinck; Consultant, CEpartner4U BV
 (name, function and signature of authorized representative)

Appendix

List of devices.

Device types	Item# Accusure® ELISA Microwells	Item# Accusure® CLIA Microwells	Item# QSure® Control	Item# Instrum ent	EDMS code	Risk Class	First date of CE-marking
Thyroid							
Total Triiodothyronine (T3) 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 (T4) 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 (T3 SBS) Test System	8125-300	8175-300			12.04.01.01.00	Low	2010-06-29
Total Thyroxine (T4 SES) 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 17OHHP (N-17OHHP) 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-Thyroperoxidase (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# OSure® Control	Item# Instum ent	EDMS code	Risk Class	First date of CE-marking
-hCG) Test System							
Human Chorionic Gonadotropin (hCG), Human Prolactin (hPL), Human Luteinizing Hormone (LH), Folic Acid Stimulating Hormone (FSH) Fertility Panel (VAST) Test System	8325-300	8375-300			12.05.01.90.00	Low	2005-09-24
Alpha-Fetoprotein (AFP), Human Chorionic Gonadotropin (hCG), Unconjugated Estriol (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 Estriol (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 - S1 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	5875-300			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# Instum 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 (βhCG) 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, Estradiol			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			IN006-2		21.02.10.01	Low	2013-09-16
Lumax CLIA Analyzer			IN001		21.02.10.01	Low	2006-06-24
Neo-Lumax CLIA Analyzer			IN010		21.02.10.01	Low	2011-09-26



Declaration of Conformity

2013-09 DoC_MB_v08

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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
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
Lumax36 CLIA Analyzer				IN004	21.02.10.01	Low	2007-09-01
LuMatic CLIA Analyzer				IN008	21.02.10.01	Low	2011-09-28
Eldex 3.8 ELISA Analyzer				IN003	21.02.10.01	Low	2007-09-10
Neo-Elex ELISA Analyzer				IN009	21.02.10.01	Low	2011-09-28
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



ERMA INC.

2-31-6 Yushima, Bunkyo-ku, Tokyo 113-0034; Japan
Phone: 81-3-3818-6281 Fax: 81-3-3813-7301
E-mail: address: trade@erma.co.jp.

MANUFACTURER'S AUTHORIZATION LETTER

Date Apr. 13, 2007

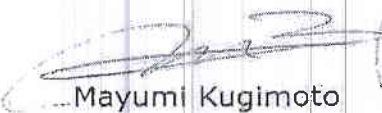
To whom it may concern,

WHEREAS, we ERMA INC., who are official and reputable manufacturers of medical and laboratory equipment as specified hereafter, hematology analyzer, biochemical analyzer, bilirubinmeter, hemoglobinmeter, microtome, having factories at 2-31-6 Yushima Bunkyo-ku, Tokyo 113-0034 Japan do hereby authorize:

GLOBAL BIOMARKETING GROUP MOLDOVA
Office 607, Tighina Str. 65 Chisinau, Moldova

To submit a bid in relation to the Bidding Document indicated above, the purpose of which is to provide the goods manufactured by us and to subsequently and sign the Contract.

Truly yours,


Mayumi Kugimoto
Manager, International Div.
ERMA INC.





CERTIFICAT CERTIFICATE

N° A 3001

Nous certifions par la présente que le Système de Management de la Qualité de la société :
We hereby certify that the Quality Management System of the company:

**BIOLABO
LES HAUTES RIVES
02160 Maizy - France**

est conforme aux exigences de la norme suivante :
is in compliance with the requirements of the following standard:

ISO 9001 : 2015

Le domaine d'application du Système de Management de la Qualité est le suivant :
The scope of the Quality Management System is:

**CONCEPTION, FABRICATION ET VENTE DE DISPOSITIFS
MÉDICAUX DE DIAGNOSTIC IN VITRO. SUPPORT
TECHNIQUE ET SERVICE D'ASSISTANCE.**

**DESIGN, MANUFACTURING AND SALE OF IN VITRO DIAGNOSTIC MEDICAL DEVICES.
TECHNICAL SUPPORT AND SUPPORT SERVICES.**

Ce certificat demeurera en vigueur jusqu'à sa fin de validité à moins d'avis contraire, à condition que la mise en place et la conformité du Système de Management de la Qualité soient jugées satisfaisantes lors des audits de surveillance et que les conditions du contrat de AB Certification soient observées.

This certificate is valid until its expiry date unless further notice, provided that the compliance and implementation of the Quality Management System are found to be satisfactory at follow-up audits and that AB Certification contract rules are fulfilled.

Fait à PARIS, le 24 décembre 2018
Signed in PARIS on the 24th of December 2018

Date de validité : 23 décembre 2021
Expiry date: 23rd of December 2021



Georges ABI RACHED
Le Représentant d'AB Certification
AB Certification Representative

Le Représentant de l'Entreprise
The Company Representative

Organisme accrédité COFRAC N° 4-0023
Accredited body by COFRAC N° 4-0023



CERTIFICAT CERTIFICATE

N° A 3001

Nous certifions par la présente que le Système de Management de la Qualité des Dispositifs Médicaux de la société :
We hereby certify that the Medical Devices Quality Management System of the company:

BIOLABO
LES HAUTES RIVES
02160 Maizy - France

est conforme aux exigences de la norme suivante :
is in compliance with the requirements of the following standard:

ISO 13485 : 2016

Le domaine d'application du Système de Management de la Qualité des Dispositifs Médicaux est le suivant :
The scope of the Medical Devices Quality Management System is as follows:

**CONCEPTION, FABRICATION ET VENTE DE DISPOSITIFS
MÉDICAUX DE DIAGNOSTIC IN VITRO. SUPPORT
TECHNIQUE ET SERVICE D'ASSISTANCE.**

*DESIGN, MANUFACTURING AND SALE OF IN VITRO DIAGNOSTIC MEDICAL DEVICES.
TECHNICAL SUPPORT AND SUPPORT SERVICES*

Ce certificat demeure en vigueur pour une période de trois ans à moins d'avis contraire, à condition que la mise en place et la conformité du Système de Management de la Qualité des Dispositifs Médicaux soient jugées satisfaisantes lors des audits de surveillance et que les conditions du contrat de AB Certification soient observées.
This certificate is valid for a three-year period unless further notice, provided that the compliance and implementation of the Medical Devices Quality Management System are found to be satisfactory at follow-up audits and that AB Certification contract rules are fulfilled.

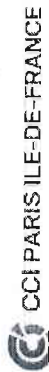
Fait à PARIS, le 24 décembre 2018
Signed in PARIS on the 24th of December 2018

Date de validité : 23 décembre 2021
Expiry date: 23rd of December 2021



Georges ABI RACHED
Le Représentant d'AB Certification
AB Certification Representative

Le Représentant de l'Entreprise
The Company Representative



Direction Générale Adjointe - Services aux Entreprises et Développement International
 Direction des réseaux et partenariats internationaux
 Service CLV

Certificat de Libre Vente pour l'exportation vers les pays non membres de l'Union Européenne
Free sale certificate for exportation to the non-EC Member States

dispositifs médicaux de diagnostic in vitro relevant de la directive n°98/79/CE
in vitro diagnostic medical devices covered by Directive 98/79/EC

PARTIE A COMPLÉTER PAR LE DEMANDEUR

Section to be completed by the applicant

Catégorie(s) du(des) dispositif(s) : : Réactifs et instruments de laboratoires pour la Biologie Médicale

Device(s) category: Reagents & Instruments for Medical Biology

Nombre de page en annexe : 5

Page in annex : 5

La désignation du(des) dispositif(s) apparaît sur la déclaration(s) CE de conformité du fabricant ou du mandataire
The name of the device(s) appears on the EC declaration(s) of conformity of the manufacturer or the authorized representative

Classification du(des) dispositif(s) :

dispositif de l'annexe II liste A

dispositif de l'annexe II

autotest hors annexe II

autre dispositif (tous les dispositifs sauf dispositifs de l'annexe II et autotests)

device for self-testing not listed in annex II
other device (all devices except annex II and self-testing devices)

Nom et adresse du fabricant ou du mandataire :
Name and address of the manufacturer or the authorized representative:

BIOLABO SAS / Mr. Jean François CHARPENTIER, Les Hautes Rives 02160 MAIZY

Nom et adresse du site de production (facultatif)

BIOLABO S.A.S, Les Hautes Rives 02160 MAIZY

Je soussigné Isabelle, Oget, Directrice Affaires Réglementaires certifie que les informations mentionnées ci-dessus sont exactes et que les dispositifs médicaux de diagnostic in vitro figurant sur la(les) déclaration(s) CE de conformité sont marqués CE sous ma responsabilité au titre de la directive n°98/79/CE et répondent aux exigences essentielles de santé et de sécurité.

I the undersigned Isabelle, Oget, Director of Regulatory Affairs declare that the information above-mentioned is correct and the in vitro diagnostic medical devices on the CE declaration(s) of conformity are CE marked under my responsibility within the meaning of the European directive n°98/79/EC and fulfil the essential requirements of health and safety.

Date : 30/08/2018

Signature :

(Handwritten signature and stamp of BIOLABO S.A.S)

PARTIE RESERVEE A LA CCIR PARIS IDF

Section reserved for the administration

Les dispositifs médicaux de diagnostic in vitro marqués CE en conformité avec la directive 98/79/CE peuvent être mis sur le marché en France et dans les autres Etats membres de l'Union Européenne et parties à l'accord sur l'espace économique européen, et être exportés vers les pays tiers. Ce certificat de libre vente est valide à concurrence du maintien, par le fabricant des dispositifs concernés, d'une déclaration de conformité (autre dispositifs), accompagnée le cas échéant, des certificats nécessaires délivrés par un organisme notifié (dispositif de l'annexe II liste A et liste B, autotests hors annexe II). Ce certificat de libre vente est utilisable uniquement à des fins d'exportation hors Union européenne.

CCIR Paris IDF / DGA-SEDI
 Le Responsable du département
 des Facilitations du Commerce Extérieur
 CCIR Paris IDF
 9, rue Coquillière
 75001 PARIS

(Official stamp of CCIR Paris IDF)

The in vitro diagnostic medical devices CE marked in conformity with the directive 98/79/EC can be placed on the French market and in the other Member States. This free sale certificate is valid until the maintenance, by the manufacturer of the concerned devices, of an CE declaration of conformity (other devices) together with when appropriate, the certificates delivered by a notified body (devices of list A and B, annex II, devices for self-testing not listed in annex II). This free sale certificate can only be used for exportation outside European Union.

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
LP80501	ACIDE URIQUE Méthode Uricase	URIC ACID Uricase Method
LP80601	ACIDE URIQUE Méthode Uricase	URIC ACID Uricase Method
80002	ALBUMINE Méthode BCG	ALBUMIN BCG Method
99029	ALCOOL Ethanol	ALCOHOL Ethanol
99059	ALCOOL Ethanol	ALCOHOL Ethanol
80327	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
LP80507	ALT TGP (IFCC)	ALT GPT (IFCC)
LP80607	ALT TGP (IFCC)	ALT GPT (IFCC)
92027	ALT TGP Méthode Colorimétrique	ALT GPT Colorimetric Method
99523	AMYLASE CNPG3	AMYLASE CNPG3
99123	AMYLASE CNPG3	AMYLASE CNPG3
LP99553	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
LP80505	AST TGO (IFCC)	AST GOT (IFCC)
LP80605	AST TGO (IFCC)	AST GOT (IFCC)
92025	AST TGO Méthode Colorimétrique	AST GOT Colorimetric Method
92026	Solution Soude 0,4 N	NPOH 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 AND 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
80108	CHOLESTEROL CHOD-PAP	CHOLESTEROL CHOD-PAP
LP80108	CHOLESTEROL CHOD-PAP	CHOLESTEROL CHOD-PAP
87656	CHOLESTEROL CHOD-PAP	CHOLESTEROL CHOD-PAP
88656	CHOLESTEROL Non estérifié CHOD-PAP	Non Esterified CHOLESTEROL CHOD-PAP
98656	CHOLESTEROL Non estérifié CHOD-PAP	Non Esterified CHOLESTEROL CHOD-PAP
87356	CHOLESTEROL CHOD-PAP	CHOLESTEROL CHOD-PAP
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
88536	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	Isenzyme CK-MB Méthode d'immunoséquestration	CK-MB Isoenzyme Immunoséquestration Method
97317	Isenzyme CK-MB Méthode d'immunoséquestration	CK-MB Isoenzyme Immunoséquestration Method

BIOLABO - Désignation des Dispositifs / Devices Designation p2/5

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) Bathophénaanthroline	IRON (SFBC) Bathophenanthroline
92108	FER Méthode directe (Féridène)	T.I.B.C. Total Iron Binding Capacity
92308	C.T.F. Capacité Totale de Fixation du Fer	U.I.B.C. Unsaturated Iron Binding Capacity
97408	C.L.F. Capacité Latente de Fixation du Fer	G6-PDH U.V. Kinetic Method
97089	G6-PDH Méthode cinétique U.V.	Lyophilised G6-PDH U.V. Kinetic Method
97099	G6-PDH lyophilisée Méthode cinétique U.V.	GAMMA GT carboxy 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
16GL8	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
LP80209	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
LP87809	GLUCOSE GOD-PAP	GLUCOSE GOD-PAP
3502200	HEMOGLOBINE Méthode Colorimétrique (Cyanmethémoglobine)	HAEMOGLOBIN Colorimetric Method (Cyanmethemoglobin)
82250	HEMOGLOBINE Méthode Colorimétrique (Cyanmethémoglobine)	HAEMOGLOBIN Colorimetric Method (Cyanmethemoglobin)
92011	L.D.H. (LDHP) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
92111	L.D.H. (LDHP) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
92511	L.D.H. (LDHP) Méthode SFBC modifiée	L.D.H. (LDH-P) SFBC Modified Method
99681	LIPASE Méthode cinétique	LIPASE Kinetic Method
99891	LIPASE Méthode cinétique	LIPASE Kinetic Method
87212	MAGNESIUM Calmagite	MAGNESIUM Calmagite
92214	MAGNESIUM CALMAGITE Haute Stabilité - Haute Linéarité	MAGNESIUM CALMAGITE High Stability - High Linearity
92214	PHOSPHATASE ALCALINE (DEA)	ALKALINE PHOSPHATASE (DEA)
82860	PHOSPHATASE ACIDE Méthode Cinétique	ACID PHOSPHATASE Kinetic Method
3300060	PHOSPHATASE ACIDE Méthode Point Final (PNFP)	ACID PHOSPHATASE End Point Method (PNFP)
99105	PHOSPHOLIPIDES Méthode colorimétrique enzymatique	PHOSPHOLIPIDS Colorimetric enzymatic Method
99110	PHOSPHORE Inorganique Méthode U.V.	PHOSPHOLIPIDS Colorimetric enzymatic Method
80015	PROTEINES TOTALES Méthode Buret	Inorganic PHOSPHORUS U.V. Method
80016	PROTEINES TOTALES Méthode Buret	TOTAL PROTEIN Buret Method
LP87016	PROTEINES U.S. Méthode Rouge de Pyrogallol	TOTAL PROTEIN Buret Method
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
LP80519	TRIGLYCERIDES Méthode GPO	TRIGLYCERIDES GPO Method
LP80619	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
LP99532	UREE U.V. Méthode Cinétique Haute Linéarité	UREA U.V. High Linearity Kinetic Method
LP99632	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 p3/5

REF	DESIGNATION FR	DESIGNATION GB
95010	BIOLABO EXATROL-N Taux 1	BIOLABO EXATROL-N Level 1
95011	BIOLABO EXATROL-P Taux 2	BIOLABO EXATROL-P Level 2
95015	BIOLABO MULTICALIBRATOR Calibrateur Multiparamétrique	BIOLABO MULTICALIBRATOR Multiparametric calibrator
95020	BIOLABO EQQ Evaluation externe de la qualité	BIOLABO EQA External Quality Assessment
95403	BIOLABO CONTROLE PEDIATRIQUE	BIOLABO PAEDIATRIC CONTROL
95406	CALIBRATEUR CHOLESTEROL-HDL	HDL-CHOLESTEROL CALIBRATOR
95906	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 Lipids Level 1
95526	Sérum de contrôle HDL LDL CK-MB Lipides Taux 2	Control serum HDL LDL CK-MB Lipids Level 2
95601	Calibrateur LIPASE	LIPASE Calibrator
95013	Contrôle Normal AMMONIAC ALCOOL BICARBONATE	Normal Control AMMONIAC ALCOOL BICARBONATE
95023	Contrôle Pathologique AMMONIAC ALCOOL BICARBONATE	Pathological Control AMMONIAC ALCOOL BICARBONATE
95089	G6-PDH Contrôle normal (hémolysat humain lyophilisé)	G6-PDH Normal control (Lyophilised human hemolysed blood)
95289	G6-PDH Contrôle Déficent (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 Taux de Prothrombine (TP)	BIO-TP Prothrombin Time (PT)
13881	BIO-TP Taux de Prothrombine (TP)	BIO-TP Prothrombin Time (PT)
13702	BIO-TP U (Low IS) Taux de Prothrombine (TP)	BIO-TP U (Low IS) Prothrombin Time (PT)
13704	BIO-TP U (Low IS) Taux de Prothrombine (TP)	BIO-TP U (Low IS) Prothrombin Time (PT)
13712	BIO-TP U (Low IS) Taux de Prothrombine (TP)	BIO-TP U (Low IS) Prothrombin Time (PT)
13883	TAMPON OWREN KOLLER	OWREN KOLLER BUFFER
13560	BIO-CK TCA Kaolin	BIO-CK APTT Kaolin
13570	BIO-CK TCA Silice	BIO-CK APTT Silica
13560	BIO-SIL TCA Silice	BIO-SIL APTT Silica
13670	BIO-SIL TCA Silice	BIO-SIL APTT Silica
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
13980	BIO-TT Temps de Thrombine	BIO-TT Thrombin Time
13565	TP-CALSET Set de Plasmas de Référence	TP-CALSET Standard Set
13970	BIO-CAL Plasma de référence	BIO-CAL Reference Plasma
13961	PLASMA CONTRÔLE Taux 1	CONTROL PLASMA Level 1
13962	PLASMA CONTRÔLE Taux 2	CONTROL PLASMA Level 2
13963	PLASMA CONTRÔLE Taux 3	CONTROL PLASMA Level 3
13210	D-DIMER Test Immunoturbidimétrique	D-DIMER Turbidimetric immunoassay
13211	D-DIMER Contrôle 1	D-DIMER Control 1
13212	D-DIMER Contrôle 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

BIOLABO - Désignation des Dispositifs / Devices Designation p4/5

BIOLABO - Désignation des Dispositifs / Devices Designation p5/5

REF	DESIGNATION FR	DESIGNATION GB	REF	DESIGNATION FR	DESIGNATION GB
13306	FACTOR VIII Plasma Déficient	FACTOR VIII Deficient plasma	SLO CALSET41	BIOLABO ASLO Kit de Calibration	BIOLABO ASLO Standard Set
13308	FACTOR IX Plasma Déficient	FACTOR IX Deficient plasma	ASLO CONT1	BIOLABO ASLO Contrôle	BIOLABO ASLO Control
13310	FACTOR X Plasma Déficient	FACTOR X Deficient plasma	ASLO CONT5	BIOLABO ASLO Contrôle	BIOLABO ASLO Control
13311	FACTOR XI Plasma Déficient	FACTOR XI Deficient plasma	APOA1620E	APOLIPOPROTEINE A1 Test Immunoturbidimétrique	APOLIPOPROTEINE A1 Turbidimetric Immunoassay
13312	FACTOR XII Plasma Déficient	FACTOR XII Deficient plasma	APOB620E	APOLIPOPROTEINE B Test Immunoturbidimétrique	APOLIPOPROTEINE B Turbidimetric Immunoassay
13971	COATROL 1 Taux 1	COATROL 1 Level 1	APOA1050E	APOLIPOPROTEINE A1 Test Immunoturbidimétrique	APOLIPOPROTEINE A1 Turbidimetric Immunoassay
13972	COATROL 2 Taux 2	COATROL 2 Level 2	APOB050E	APOLIPOPROTEINE B Test Immunoturbidimétrique	APOLIPOPROTEINE B Turbidimetric Immunoassay
9905TH	S. Typhi H (g-H)	S. Typhi H (g-H)	BIOLABO A1B Calibrant Haut	BIOLABO A1B Calibrant Haut	BIOLABO A1B Standard High
9905TO	S. Typhi O (9,12-O)	S. Typhi O (9,12-O)	BIOLABO A1B Contrôle	BIOLABO A1B Contrôle	BIOLABO A1B Control
9905AH	S. Paratyphi AH (e-H)	S. Paratyphi AH (e-H)	MICROALBUMINE Test Immunoturbidimétrique	MICROALBUMINE Test Immunoturbidimétrique	MICROALBUMIN Turbidimetric Immunoassay
9905AO	S. Paratyphi AO (1,2,12-O)	S. Paratyphi AO (1,2,12-O)	MICROALBUMINE Test Immunoturbidimétrique	MICROALBUMINE Test Immunoturbidimétrique	MICROALBUMIN Turbidimetric Immunoassay
9905BH	S. Paratyphi BH (b-H)	S. Paratyphi BH (b-H)	MICROALBUMINE Calibrant Super Haut	MICROALBUMINE Calibrant Super Haut	MICROALBUMIN Standard Super High
9905BO	S. Paratyphi BO (1,4,5-O)	S. Paratyphi BO (1,4,5-O)	MICROALBUMINE Kit de calibration	MICROALBUMINE Kit de calibration	MICROALBUMIN Standard Set
9905CH	S. Paratyphi CH (c-H)	S. Paratyphi CH (c-H)	MICROALBUMINE Contrôle	MICROALBUMINE Contrôle	MICROALBUMIN Control
9905CO	S. Paratyphi CO (6,7-O)	S. Paratyphi CO (6,7-O)	HbA1c ENZYM	HbA1c ENZYM	HbA1c ENZYM
9905EA	Brucella abortus	Brucella Abortus	HbA1c ENZYM Kit de calibration	HbA1c ENZYM Kit de calibration	HbA1c ENZYM Standard Set
9905FA	Proteus OXK	Proteus OXK	HbA1c Test Immunoturbidimétrique	HbA1c Test Immunoturbidimétrique	HbA1c Turbidimetric Immunoassay
9905FP	Proteus OX19	Proteus OX19	HbA1c Test Immunoturbidimétrique	HbA1c Test Immunoturbidimétrique	HbA1c Turbidimetric Immunoassay
9905P2	Proteus OX2	Proteus OX2	HbA1c Kit de calibration	HbA1c Kit de calibration	HbA1c Standard Set
9905BM	Brucella Melitensis	Brucella Melitensis	HbA1c Kit de contrôle	HbA1c Kit de contrôle	HbA1c Control Set
9905RB	Rose Bengal (B. Abortus)	Rose Bengal (B. Abortus)	KENZA MAX BioChemistry PHOTOMETRE	KENZA MAX BioChemistry PHOTOMETRE	KENZA MAX BioChemistry PHOTOMETRE
9905PC	Contrôle Positif Polyvalent	Positive Polyvalent Control	KENZA 120TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 120TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 120TX - AUTOMATIC BIOCHEMISTRY ANALYSER
9905NC	Contrôle Négatif Polyvalent	Negative Polyvalent Control	KENZA 240TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 240TX - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE	KENZA 240TX - AUTOMATIC BIOCHEMISTRY ANALYSER
9905F4	ANTIGENES FEBRILES Pour Tests de Widal Félix	STAINED FEBRILE ANTIGENS For Widal Felix Tests	KENZA 240ISE - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE avec module ISE	KENZA 240ISE - ANALYSEUR AUTOMATIQUE DE BIOCHIMIE avec module ISE	KENZA 240ISE - AUTOMATIC BIOCHEMISTRY ANALYSER with ISE Module
9905F6	ANTIGENES FEBRILES Pour Tests de Widal Félix	STAINED FEBRILE ANTIGENS For Widal Felix Tests	KENZA 450TX	KENZA 450TX	KENZA 450TX
9905F8	ANTIGENES FEBRILES Pour Tests de Widal Félix	STAINED FEBRILE ANTIGENS For Widal Felix Tests	KENZA 450ISE	KENZA 450ISE	KENZA 450ISE
081050	ASLO-LATEX	ASLO-LATEX	BIO SOLEA 2 - COAGULOMETRE 2 CANAUX	BIO SOLEA 2 - COAGULOMETRE 2 CANAUX	BIO SOLEA 2 - COAGULOMETER 2 CHANNELS
097100	CRP-LATEX	CRP-LATEX	BIO SOLEA 4 - COAGULOMETRE 4 CANAUX	BIO SOLEA 4 - COAGULOMETRE 4 CANAUX	BIO SOLEA 4 - COAGULOMETER 4 CHANNELS
098100	FR-LATEX	FR-LATEX	SOLEA 100 - ANALYSEUR AUTOMATIQUE D'HEMOSTASE	SOLEA 100 - ANALYSEUR AUTOMATIQUE D'HEMOSTASE	SOLEA 100 - FULL-AUTOMATED COAGULATION ANALYSER
3800100	RPR-CHARBON	RPR-CHARBON	SCUP120 Serum Cup K120TX	SCUP120 Serum Cup K120TX	Serum Cup K120TX
3800150	RPR-CHARBON	RPR-CHARBON	SERUM CUPS	SERUM CUPS	SERUM CUPS
4500100	TPHA	TPHA	CO4015 Extra Cleaning	CO4015 Extra Cleaning	Extra Cleaning
4500200	TPHA	TPHA	CO4020 Ipo Cleaning	CO4020 Ipo Cleaning	Ipo Cleaning
085100	HCG-LATEX	HCG-LATEX	CO0058 SERUM CUPS K450	CO0058 SERUM CUPS K450	SERUM CUPS K450
RF050E	Facteurs Rhumatoïdes (FR) Test Immunoturbidimétrique	RHEUMATOID FACTOR (RF) Turbidimetric Immunoassay	K450CS Cleaning Solution K450	K450CS Cleaning Solution K450	Cleaning Solution K450
RF520E	Facteurs Rhumatoïdes (FR) Test Immunoturbidimétrique	RHEUMATOID FACTOR (RF) Turbidimetric Immunoassay	RP240ISE Pack Reactifs - ISE	RP240ISE Pack Reactifs - ISE	Reagent Pack - ISE
RF CALSET51	BIOLABO FR Kit de Calibration	BIOLABO RF Standard Set	G2058FA Cleaning Solution - ISE	G2058FA Cleaning Solution - ISE	Cleaning Solution - ISE
RF CALLSH1	BIOLABO FR Calibrant Super Haut	BIOLABO RF Standard Super High	5202 Electrode K - ISE	5202 Electrode K - ISE	Electrode K - ISE
RF CONT1	BIOLABO FR Contrôle	BIOLABO RF Control	5205 Electrode LI - ISE	5205 Electrode LI - ISE	Electrode Li - ISE
RF CONT5	BIOLABO FR Contrôle	BIOLABO RF Control	5207 Electrode CI - ISE	5207 Electrode CI - ISE	Electrode C - ISE
CRP050E	CRP Test Immunoturbidimétrique	CRP Turbidimetric Immunoassay	5201 Electrode Na - ISE	5201 Electrode Na - ISE	Electrode Na - ISE
CRP620E	CRP Test Immunoturbidimétrique	CRP Turbidimetric Immunoassay	5204 Electrode de référence	5204 Electrode de référence	Reference Electrode
CRP CALSET51	BIOLABO CRP Kit de Calibration	BIOLABO CRP Standard Set	S100CS CLEANING SOLUTION SOLEA 100	S100CS CLEANING SOLUTION SOLEA 100	CLEANING SOLUTION SOLEA 100
CRP CALLSH1	BIOLABO CRP Calibrant Super Haut	BIOLABO CRP Standard Super High			
CRP CONTL1	BIOLABO CRP Contrôle Bas	BIOLABO CRP Control Low			
CRP CONTL5	BIOLABO CRP Contrôle Bas	BIOLABO CRP Control Low			
CRP CONTH1	BIOLABO CRP Contrôle Haut	BIOLABO CRP Control High			
CRP CONTH5	BIOLABO CRP Contrôle Haut	BIOLABO CRP Control High			
ASLO050E	ASLO Test Immunoturbidimétrique	ASLO Turbidimetric Immunoassay			
ASLO620E	ASLO Test Immunoturbidimétrique	ASLO Turbidimetric Immunoassay			
ASLO CALH1	BIOLABO ASLO Calibrant Haut	BIOLABO ASLO Standard High			
ASLO CALSH1	BIOLABO ASLO Calibrant Super Haut	BIOLABO ASLO Standard Super High			

BIOLABO



Всем заинтересованным лицам

Авторизационное письмо

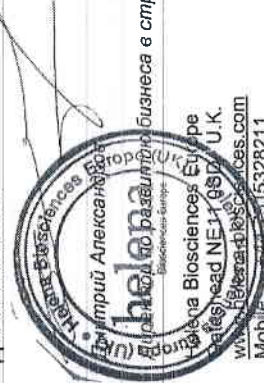
Настоящим, мы, компания «HELENA LABORATORIES (UK) Ltd», торгующая как «HELENA BIOSCIENCES EUROPE», с центральным офисом по адресу: Queensway South, Team Valley Trading Estate, Gateshead, Tyne & Wear, NE11 0SD, Великобритания, подтверждает, что:

Компания "GBG-MLD" SRL, республика Молдова, г. Кишинёв, MD-2001, улица Chisinau Tigina, дом 65, офис 607 являются уполномоченными дистрибьюторами всей продукции компании «HELENA BIOSCIENCES EUROPE» на территории Республики Молдова и авторизована принимать участие во всех тендерах.

Компани "GBG-MLD" SRL имеет право импорта, продвижения и продажи выше перечисленной продукции на территории Республики Молдова.

Настоящее письмо действительно до 31 декабря 2020 года.

Дата: 22/01/2020



Helena Biosciences Europe, Импорт Александру Бурлаку по развитию бизнеса в странах СНГ, Европы и Азии.

Helena Biosciences Europe
Gateshead NE11 0SD, U.K.
www.helena-biosciences.com
Mobile: +44 191 482 8442
Phone: +44 191 482 8462
Email: da@helena-biosciences.com

bsi.



By Royal Charter

Certificate of Registration

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

This is to certify that:

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

Holds Certificate Number:

MD 69326

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

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

For and on behalf of BSI:

Stewart Brain, Head of Compliance & Risk - Medical Devices

Original Registration Date: 2002-10-25

Latest Revision Date: 2018-11-28

Effective Date: 2018-04-14

Expiry Date: 2021-04-13

Page: 1 of 2



...making excellence a habit.™

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract.
An electronic certificate can be authenticated [online](#).
Printed copies can be validated at www.bsigroup.com/ClientDirectory

Information and Contact: BSI, Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes MK5 8PP. Tel: + 44 345 080 9000
BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK.
A Member of the BSI Group of Companies

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	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

Tel +44 (0)191 482 8440
Fax +44 (0)191 482 8442
info@helena-biosciences.com
www.helena-biosciences.com

Helena Biosciences Europe
Queensway South, Team Valley Trading Estate,
Gateshead, Tyne and Wear, NE11 0SD,
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
5559SLQ	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

Tel +44 (0)191 482 8440
Fax +44 (0)191 482 8442
info@helena-biosciences.com
www.helena-biosciences.com

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



articoli per laboratorio analisi
disposable labware

www.kima.it



Messrs

"GBG-MLD" SRL
STR. TIGHINA 65
2001 CHISINAU
MOLDOVA

Piove di Sacco, 25/02/2019

DISTRIBUTOR AGREEMENT

To whom it may concern, we hereby declare that:

KIMA sas – Via Leonardo Da Vinci 22 – 35028 piove di Sacco - (PD) - ITALY

appoints "GBG-MLD" SRL – STR. TIGHINA 65. - 2001 CHISINAU –MOLDOVA

as authorized distributor of KIMA plastic labware products in the territory of MOLDOVA

GBG MLD has the right to import and distribute KIMA plastic labware products.

This Agreement is valid one (2) years from the present date.

The Distributor does not have any possibility to oblige the company KIMA sas with quantities or delivery time as well as prices without prior written authorization from KIMA sas.

KIMA sas keeps the right to modify the prices according to the market of the raw materials:

Renzo Chiarin
Managing Director

KIMA S.R.L.
Via Leonardo Da Vinci, 22
35028 PIOVE DI SACCO (PD)
Partita IVA 01466290283



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www.iqnet-certification.com

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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,
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° 004A

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Signatory of EA, IAF and ILAC Mutual Recognition Agreements



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CERTIFICATE No. _____

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

MEUS S.r.l.

Unità Operative / Operative Units

Via Leonardo Da Vinci, 24B-26-28 - Zona Industriale Tognana - 35028 Piove di Sacco (PD) - Italia
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 stampi per articoli in plastica per laboratorio analisi.

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

Progettazione e produzione di aghi e dispositivi sterili per il prelievo ematico.

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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

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. Progettazione e produzione di stampi per articoli in plastica per laboratorio analisi.

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. Design and production of moulds for plastic labware.

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.

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17/01/2022


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CERTIFICATO n. 4265/4/B
CERTIFICATE No. _____

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

ROLL S.R.L.

UNITÀ OPERATIVE / OPERATIVE UNITS

Via Leonardo Da Vinci, 24A - 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: 14

Progettazione e produzione di Holders (camicie) per prelievo sottovuoto.
Progettazione e produzione di kit diagnostici per l'analisi del sangue e dei liquidi biologici.
Stampaggio di materie termoplastiche ad iniezione per articoli medicali.

*Design and production of Holders for vacuum sampling.
Design and production of diagnostic kits for blood and biological liquids analysis.
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.

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CERTIFICATO n. 4265/4/D
CERTIFICATE No. _____

SI CERTIFICA CHE IL SISTEMA DI GESTIONE PER LA QUALITÀ DI
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.
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.

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CISQ is the Italian Federation of management system Certification Bodies.

TO WHOM IT MAY CONCERN

Letter of Authorization

We APTACA SPA , with head offices and plant located in :

Regione Monforte nr 30

14053 Canelli (At) Italy

Confirm that the below Company :

"GBG-MLD" S.R.L.
Tighina str.65, office 607
MD-2001, Chisinau,
Republic of Moldova

Web: www.gbg.md
Ph. +373 22 54 91 20
+373 22 54 91 21

Is authorized to prepare price quotations, advertising activities, warranty service, offers, to participate in tenders and to sell our whole range of product on exclusive basis in the territory of MOLDAVIA. This letter is valid until 31/12/2020 and may be prolonged by mutual agreement.

ON BEHALF OF NUOVA APTACA S.R.L.

Veronica FERRARI

Export Manager


APTACA SPA
Reg. Monforte n. 30
Tel. 0141/835075 r.a. - Fax 0141/835292
14053 CANELLI (AT)
C.F. 07520900155 - P.I. 00862050960
Cod. Univoco: SUBM70N

Canelli 25/11/2019

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.

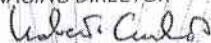
Commercializzazione di articoli da laboratorio

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
SCA N° 020D ISP N° 075E
PRS N° 097C

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