

BC "MOLDINDCONBANK" S.A. Filiala "Invest"

Republica Moldova, MD-2068 mun. Chişînău, bd. Moscovei, 14/1 Tel. : (373-22) 43-44-81, 43-46-24 Fax : (373-22) 43-44-22 cod: MOLDMD2X329

IAN. 2016 Data

Республика Молдова, MD-2068 мун. Кишинэу, бул. Московей, 14/1 Тел. : (373-22) 43-44-81, 43-46-24 Факс : (373-22) 43-44-22 код: MOLDMD2X329

Filiala "Invest" BC "Moldindconbank" SA confirmă existența contului curent in moneda nationala al "BIOSISTEM MLD" S.R.L. (c/f 1010600028048), cu IBAN MD95ML00000002251429243.

Codul băncii MOLDMD2X329.

Director

Nina **Turcan**



1 Balmiy

Nina Balmuş

Ex. Diana Brinza Tel. 43-45-96



MOLDOVA

CERTIFICAT DE ÎMREGISTRARE

Societatea cu Răspundere Limitată "BIOSISTEM MLD" ESTE ÎNREGISTRATĂ LA CAMERA ÎNREGISTRĂRII DE STAT

Numărul de identificare de stat - codul fiscal 1010600028048

Data înregistrării

Data eliberării

12.08.2010

12.08.2010

Svirepova Ludmila, registrator

Funcția, numele, prenumele persoanei care a eliberat certificatul

comnătura



MD 0101250

"CAMERA ÎNREGISTRĂRII DE STAT" Î.S. Secția fonduri speciale și informații curente

EXTRAS

din Registrul de stat al persoanelor juridice

nr. 14419 din 11.07.2016

Denumirea completă: Societatea cu Răspundere Limitată «BIOSISTEM MLD». Denumirea prescurtată: «BIOSISTEM MLD» S.R.L. Forma juridică de organizare: Societate cu Răspundere Limitată. Numărul de identificare de stat și codul fiscal: 1010600028048. Data înregistrării de stat: 12.08.2010. Sediul: MD-2001, str. Albișoara, 16/1, ap.(of.) 7, mun. Chișinău, Republica Moldova. Modul de constituire: nou creată. Obiectul principal de activitate: 1 Activitatea farmaceutică; 2 Importul, fabricarea, comercializarea, asistența tehnică și (sau) reparația dispozitivelor medicale și (sau) a opticii; 3 Acordarea asistenței medicale de către instituțiile medico-sanitare private; 4 Comerțul cu ridicata al calculatoarelor, echipamentelor periferice și software-ului; 5 Întreținerea și repararea mașinilor de birou și a tehnicii de calcul; 6 Consultații în domeniul sistemelor de calcul. Capitalul social: 5400 lei. Administrator: POIATA VITALIE, IDNP 0983103892591, Asociați: 1. POIATA VITALIE, IDNP 0983103892591 cota 1803.60 lei, ce constituie 33,4 % 2. NASEDCHIN ALEXANDR, IDNP 2002001070747 cota 1798.20 lei, ce constituie 33,3 % 3. KOJEVNIKOV DMITRII, IDNP 0972305012362 cota 1798.20 lei, ce constituie 33,3 %.

Prezentul extras este eliberat în temeiul art. 34 al Legii nr. 220-XVI din 19 octombrie 2007 privind înregistrarea de stat a persoanelor juridice și a întreprinzătorilor individuali și confirmă datele din Registrul de stat la data de: 11.07.2016.

Specialist principal tel. 022-266-252	Muinun	Lazari Aliona
	A A 7	* E A * 0 3 7 0 4 3 1 E



c/f 1010600028048; adresa: or. Chişinău, str. Albişoara 16/1 of.7 tel.+373-22-808-517, +373-22-808719, fax: +373-22-808-519. Web: www.biosistem-mld.com; e-mail: biosistem.mld@gmail.com

Lista fondatorilor Biosistem-mld SRL

Nr.	Nume, Prenume	IDNP
1.	Vitalie Poiata	0983103892591
2.	Alexandru Nasedchin	2002001070747
3.	Dmitrii Kojevnikov	0972305012362

Anexa nr.7.2 la Instrucțiunea aprobată prin ordinul IFPS nr. 400 din 14 martie 2014

CC 04 AE

CERTIFICAT

din

OT

privind lipsa sau existența restanțelor față de bugetul public național

20.01.2021

100779	0779
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1. Destinația / Назначение

Pentru participarea la proceduri de achizitii publice

2. Date despre contribuabil / Информация о налогоплательщике

Denumirea Наименование		Codul fiscal / Numărul de identificare Фискальный код / Идентификационный номер
BIOSISTEM MLD S.R.L.		1010600028048
Adresa sediului de bază (strada, numărul) Адрес основного месторасположения (улица, номер)		enumirea localității менование населенного пункта
Albisoara nr.16 bl.1 of.7	0150-SE	C.RISCANI

3. Atestarea lipsei sau existenței restanțelor conform datelor Sistemului Informațional Automatizat/ Подтверждение отсутствия или наличия недоимки согласно данных Информационной автоматизированной системы

La data emiterii prezentului certificat restanța față de bugetul public național constituie/ На дату выдачи данной справки недоимка перед национальным публичным бюджетом составляет: **0,00** lei/лей.

4. Valabil pînă la / Действителен до 04.02.2021

5. Autentificarea Serviciului Fiscal de Stat / Подтверждение Государственной налоговой службы

Sef DDF Rîşcani a DGAF mun. Chişinau rivan de tin the second second	Viorica CĂUȘ
Funcția/Должность	Numele și prenumele/Фамилия и имя
L.Ş/ M.Π. Executor: Claudia GOJAN	
104 006601001181	

Este extras din Sistemul Informațional al SFS SIA "Contul curent al contribuabilului"// 20.01.2021 ora 15:30:49 cu aplicarea prevederilor pct. 82-83 Ordin IFPS nr.400 din 14.03.2014 (Monitorul Oficial 72-77/399, 28.03.2014) NOTA (1,16)



EC DECLARATION OF CONFORMITY

BioSystems S.A., a company placed in Costa Brava 30, 08030 Barcelona (Spain) dedicated to the design, development and manufacturing of in vitro diagnostic medical devices.

Hereby DECLARES

That the products stated in the annex of five (5) pages joined herewith, meet the applicable provisions of the

Directive on in Vitro Diagnostic Medical Devices (98/79/EC)

under the specifications declared by BioSystems S.A.

It means that the products:

- complies with all applicable Essential Requirements as set out in the Annex I, and its technical documentation is performed following the requirements of the Annex III
- is classified as Other Device (all devices except Annex II and Self-Testing Devices), that is why the Conformity Assessment follows the procedure stated in the Annex III of the Directive without the intervention of a Notified Body.

Barcelona, November 6th, 2012

08030 BC

Dr. Antonio Elduque Managing director BioSystems S.A.



www.biosystems.es



CLINICAL CHEMISTRY – BIOCHEMISTRY:

a-Amylase-Direct a-Amylase-EPS a-Amvlase-Pancreatic Acid Phosphatase (ACP) Alanine Aminotransferase (ALT/GPT) Albumin Alkaline Phosphatase (ALP)-AMP Alkaline Phosphatase (ALP)-DEA AspartateAminotranferase (AST/GOT) Bilirubin (direct) Bilirubin (total and direct) Bilirubin (total) Calcium – Arsenazo Calcium – MTB Cholesterol Cholesterol HDL Cholesterol HDL direct Cholesterol HDL Precipitating reagent Cholesterol LDL direct Cholesterol LDL Precipitating reagent Cholinesterase (CHE) Citrate

Creatine Kinase (CK) Creatine Kinase-MB (CK-MB) Creatinine Fructosamine Fructose g-Glutamyltransferase (g-GT) Glucose Iron – Chromazurol Iron – Ferrozine Iron Binding Capacity Lactate Dehydrogenase (LDH) Lactate Dehydrogenase (LDH) - IFCC Lipase Magnesium Phosphorus Protein (total) Protein (urine) Pyridoxal Phosphate Triglycerides Urea/BUN-Color Urea/BUN-UV Uric Acid

CLINICAL CHEMISTRY – TURBIDIMETRY:

a1-acid Glycoprotein Albumin (Microalbuminuria) Anti-Streptolysin O (ASO) Antithrombin III Apolipoprotein A-I (Apo A-I) Apolipoprotein B (Apo B) b2-Microglobulin Complement Component C3 Complement Component C4

C-Reactive Protein (CRP) C-Reactive Protein-hs (CRP-hs) Ferritin Immunoglobulin A (IgA) Immunoglobulin G (IgG) Immunoglobulin M (IgM) Prealbumin Rheumatoid Factors (RF) Transferrin

CLINICAL CHEMISTRY – MICROCOLUMN CHROMATOGRAPHY:

17-Hydroxycorticosteroids
17-Ketosteroids
5-Aminolevulinic Acid (ALA) / Porphobilinogen (PBG)
5-Hydroxyindoleacetic acid (5-HIAA) Hemoglobin A1C Hemoglobin A2 Metanephrines Vanilmandelic Acid



CLINICAL CHEMISTRY – STANDARDS and CALIBRATORS:

a-1-acid Glycoprotein Standard Adenosine Deaminase (ADA) Standard Albumin (Microalbuminuria) Standard Anti-Streptolysin O (ASO) Standard Antithrombin III Standard Apolipoprotein A-I Standard Apolipoprotein B Standard b2-Microglobulin Standard Bilirubin Standard Biochemistry Calibrator Biochemistry Calibrator (Human) Cholesterol HDL/LDL Calibrator CRP/CRP-hs Standard Ferritin Standard Hemoglobin A1C-Turbi (HbA1C-Turbi) Standard Prealbumin Standard Protein Calibrators Protein (urine) Standard Rheumatoid Factors (RF) Standard

CLINICAL CHEMISTRY - INSTRUMENTS:

A15 A25 BA400 BTS-350

CLINICAL CHEMISTRY – BIOCHEMISTRY – REAGENTS AUTOMATED SYSTEMS:

a-Amylase-Direct a-Amylase-Pancreatic Adenosine Deaminase (ADA) Alanine Aminotransferase (ALT/GPT) Albumin Alkaline Phosphatase (ALP)-AMP Alkaline Phosphatase (ALP)-DEA Aspartate Aminotransferase (AST/GOT) Bilirubin (direct) Bilirubin (total) Calcium-Arsenazo Cholesterol Cholesterol HDL direct Cholesterol LDL direct Creatine Kinase (CK) Creatine Kinase-MB (CK-MB) Creatinine g-Glutamyltransferase (g-GT) Glucose Iron Ferrozine Lactate dehydrogenase (LDH) Lipase Magnesium Phosphorus Protein (total) Protein (urine) Triglycerides Urea/BUN UV Uric acid



CLINICAL CHEMISTRY – TURBIDIMETRY – REAGENTS AUTOMATED SYSTEMS:

Albumin (Microalbuminuria) Anti-Streptolysin O (ASO) Antithrombin III Complement Component C3 Complement Component C4 C-Reactive Protein (CRP) C-Reactive Protein-hs (CRP-hs) Ferritin Hemoglobin A1C-Turbi (HbA1C-Turbi) Immunoglobulin A (IgA) Immunoglobulin G (IgG) Immunoglobulin M (IgM) Rheumatoid Factors (RF) Transferrin

CLINICAL CHEMISTRY - INTERNAL QUALITY CONTROL:

ADA Controls Biochemistry Control Serum (Human) I Biochemistry Control Serum (Human) II Biochemistry Control Serum I Biochemistry Control Serum II CK-MB Control Serum Control Urine Fertility Biochemistry Control Hemoglobin A1C Control (Elevated)

Hemoglobin A1C Control (Normal) Hemoglobin A2 Control Lipid Control Serum I Lipid Control Serum II Protein Control Serum I Protein Control Serum II Rheumatoid Control Serum I Rheumatoid Control Serum II

AUTOIMMUNITY – IFA (IMMUNOFLUORESCENCE):

Anti-Adrenal Cortex Antibodies (AACA) Anti-Endomysium Antibodies (AEA) Anti-Islet Cell Antibodies (AICA) Anti-Islet Cell Antibodies (AICA) Anti-Mitochondrial Antibodies (AMA) Anti-nDNA antibodies (nDNA) Anti-Neutrophil Cytoplasmic Antibodies (ANCA) Anti-Nuclear Antibodies HEp-2 (ANA HEp-2) Anti-Nuclear Antibodies RL (ANA-RL) Anti-Skin Antibodies (ASA) Anti-Smooth Muscle Antibodies (ASMA) Anti-Striated Muscle Antibodies (AStMA)

Anti-Thyroid Antibodies (ATA) Autoantibodies DUO-HEp2/ML (DUO-HEp2/ML) Autoantibodies MsK/MsS (AA-MsK/MsS) Autoantibodies MsL/MsK/MsS (AA-MsL/MsK/MsS) Autoantibodies RK/RS (AA-RK/RS) Autoantibodies RL/RK/RS (AA-RL/RK/RS) Autoantibodies RL/RKm/RS (AA-RL/RKm/RS) Glomerular Basement Membrane Antibodies (GBMA)



AUTOIMMUNITY - ELISA:

ANA Screening Anti-Annexin V IgG/IgM (ANX) Anti-b2-Glycoprotein 1 IaG/IaM (b2GP1) Anti-Cardiolipin Antibodies (ACA-IaG/IaM) Anti-Centromere B Antibodies (CENP-B) Anti-Citrullinated Protein Antibodies (ACPA) Anti-Deamidated Gliadin Peptides IgA (DGP IgA) Anti-Deamidated Gliadin Peptides IgG (DGP IgG) Anti-dsDNA Antibodies Anti-GBM Antibodies - EIA (GBM) Anti-Gliadin Antibodies (AGA-IgG/IgA) Anti-Histones Antibodies (HIST) Anti-Insulin Antibodies (INS) Anti-Jo1 Antibodies Anti-M2 Antibodies (M2)

Anti-MPO Antibodies Anti-Nucleosome Antibodies (NCL) Anti-Phospholipid IgG/IgM (APLA) Anti-PR3 Antibodies Anti-Ribosomal P Antibodies (Rib P) Anti-Scl70 Antibodies Anti-Sm Antibodies Anti-Sm/RNP Antibodies Anti-SSA (Ro) Antibodies Anti-SSB (La) Antibodies Anti-Thyroglobulin Antibodies (Anti-Tg) Anti-Thyroid Peroxidase Antibodies (Anti-TPO) Anti-tTransglutaminase IgA Antibodies (Anti- tTG IgA) Anti-tTransglutaminase IgG Antibodies (Anti- tTG IgG) ASCA-IgG/IgA (ASCA) **ENA 4-Profile ENA 6-Screening**

AUTOINMUNIDAD – INSTRUMENTOS: AUTOIMMUNITY – INSTRUMENTS:

iPRO

RAPID TESTS – LATEX AGGLUTINATION:

Anti-Streptolysin O (ASO) - Slide C-Reactive Protein (CRP) - Slide Rheumatoid factors (RF) - Slide

INFECTIOUS IMMUNOLOGY – SYPHILIS:

RPR-Carbon

TPHA

INFECTIOUS IMMUNOLOGY – FEBRILE ANTIGENS:

Febrile Serodiagnostics Multiscreening Febrile Serodiagnostics Salmonella Brucella abortus Brucella abortus, Rose Bengal Proteus Ox19 Salmonella paratyphi AH Salmonella paratyphi AO Salmonella paratyphi BH Salmonella paratyphi BO Salmonella paratyphi CH Salmonella paratyphi CO Salmonella typhi H Salmonella typhi O Brucella Positive Control **Proteus Positive Control** Salmonella Positive Control Serology Negative Control

Certificate

Standard

ISO 9001:2015

Certificate Registr. No.

01 100 6696

Certificate Holder:

BIOSY STEMS S.A. Costa Brava 30 08030 Barcelona Spain

Scope:

Design, development, manufacture, distribution, servicing of: -Instruments and reagents for clinical diagnostic. -Instruments and reagents for agro-alimentary analysis. Distribution and service of reagents and instruments for veterinary diagnosis.

Proof has been furnished by means of an audit that the requirements of ISO 9001:2015 are met.

Validity:

The certificate is valid from 2019-12-19 until 2022-12-18. First certification 1996

2019-12-20

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln



www.tuv.com





Annex to certificate

Standard

ISO 9001:2015

Certificate Registr. No. 01 100 6696

No.

Location

Scope

/02 BIOSYSTEMS S.A. Pol. Ind. Can Tapioles naus 7-12-13 08110 Montcada i Reixac Spain

Labeling and assembly of reagent. Storage, and shipping of: - Instruments and reagents for diagnosis and reagents for clinical diagnosis.- Instruments and reagents for agrifood analysis.- Instruments and reagents for veterinary diagnosis.

2019-12-20



TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln

Page 1 of 1



Klicken Sie hier, um Text einzugeben.

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Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

BIOSYSTEMS S.A. Costa Brava 30 08030 Barcelona Spain

has established and applies a quality management system for medical devices for the following scope:

Design and development, manufacture, distribution and servicing of instruments and reagents for clinical diagnostic (see attachment for sites included)

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance.

Effective Date:

2020-01-08

Certificate Registration No.: SX 60145545 0001

An audit was performed. Report No.: 28300434 004

This Certificate is valid until: 2022-12-12

DAKKS Deutsche Akkreditierungsstelle D-ZM-14169-01-02

Turnhainland Turnhainland

Certification Body

D. Swiatko

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg Tel.: +49 221 806-1371 Fax: +49 221 806-3935 e-mail:cert-validity@de.tuv.com http://www.tuv.com/safety

Date 2020-01-08



Doc. 1/1, Rev. 0

TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment to Certificate Registration No.: Report No.:

SX 60145545 0001 28300434 004

Organization:

BIOSYSTEMS S.A. Costa Brava 30 08030 Barcelona Spain

Scope:

Site included:

Polígono Industrial Can Tapioles Naves 7, 12 y 13 08110 Montcada i Reixac Spain

Activity: Labelling and assembling of reagents, warehousing and shipment of instruments and reagents for clinical diagnostic



11/020 h 04.08 @ TÜV, TUEV and TUV are registered trademarks. Utilisation and application requires prior approval.

Date: 2020-01-08

Certification Body



D. Swiatko

CE Declaration of Conformity CE

According to the In Vitro Diagnostic Medical Devices Directive 98/79/EC

Manufacturer:	Dirui Industrial Co., Ltd.		
	95 Yunhe Street New& High Tech. Development Zone		
	Changchun Jilin 130012 P.R. China		
Authorized	Emergo Europe		
Representative :	Molenstraat 15 2513 BH The Hague		
	The Netherlands		
Medical	Product Name: Reagent strips for Urinalysis		
Device :	IVDD-Classification: Professional use		
	Lot/batches/Serial mber, Type, Periods of manufacture (where applicable)		
	DIRUI 1 ITEMS (GLU) DIRUI 1 ITEMS (KET) DIRUI 1 ITEMS (PRO) DIRUI 2 ITEMS (PRO,GLU) DIRUI 2 ITEMS(KET,GLU) DIRUI 3 ITEMS(PRO,PH,GLU) DIRUI 3 ITEMS (PRO, KET,GLU) DIRUI 4 ITEMS (PRO,PH.BLD,GLU) DIRUI 4 ITEMS (PRO,PH,SG,GLU) DIRUI 5 ITEMS (PRO,PH, BLD,KET,GLU) DIRUI 8 ITEMS DIRUI H8 DIRUI 9 ITEMS		
C C C C C C C	DIRUI 2 ITEMS (PRO,GLU)DIRUI 2 ITEMS(KET,GLU)DIRUI 3 ITEMS(PRO,PH,GLU)DIRUI 3 ITEMS (PRO, KET,GLU)DIRUI 4 ITEMS (PRO,PH.BLD,GLU)DIRUI 4 ITEMS (PRO,PH,SG,GLU)DIRUI 5 ITEMS (PRO,PH, BLD,KET,GLU)DIRUI 8 ITEMSDIRUI 8 ITEMSDIRUI H8		
	DIRUI 2 ITEMS (PRO,GLU) DIRUI 2 ITEMS(KET,GLU) DIRUI 3 ITEMS(PRO,PH,GLU) DIRUI 3 ITEMS (PRO, KET,GLU) DIRUI 4 ITEMS (PRO,PH.BLD,GLU) DIRUI 4 ITEMS (PRO,PH,SG,GLU) DIRUI 5 ITEMS (PRO,PH, BLD,KET,GLU) DIRUI 8 ITEMS DIRUI H8 DIRUI 9 ITEMS DIRUI 410 DIRUI H10 DIRUI E10 DIRUI M10 DIRUI H10-800 DIRUI H11 DIRUI H11-MA DIRUI H11-800		
	DIRUI 2 ITEMS (PRO,GLU) DIRUI 2 ITEMS(KET,GLU) DIRUI 3 ITEMS(PRO,PH,GLU) DIRUI 3 ITEMS (PRO, KET,GLU) DIRUI 4 ITEMS (PRO,PH.BLD,GLU) DIRUI 4 ITEMS (PRO,PH,SG,GLU) DIRUI 5 ITEMS (PRO,PH, BLD,KET,GLU) DIRUI 8 ITEMS DIRUI H8 DIRUI 9 ITEMS DIRUI 9 ITEMS DIRUI A10 DIRUI H10 DIRUI E10 DIRUI M10 DIRUI H10-800 DIRUI H11 DIRUI H11-MA DIRUI H11-800 DIRUI H11 A00MA DIRUI H12-800MA		
	DIRUI 2 ITEMS (PRO,GLU) DIRUI 2 ITEMS(KET,GLU) DIRUI 3 ITEMS(PRO,PH,GLU) DIRUI 3 ITEMS (PRO, KET,GLU) DIRUI 4 ITEMS (PRO,PH.BLD,GLU) DIRUI 4 ITEMS (PRO,PH,SG,GLU) DIRUI 5 ITEMS (PRO,PH, BLD,KET,GLU) DIRUI 8 ITEMS DIRUI H8 DIRUI 9 ITEMS DIRUI 410 DIRUI H10 DIRUI E10 DIRUI M10 DIRUI H10-800 DIRUI H11 DIRUI H11-MA DIRUI H11-800		

The undersigned hereby declares that the In Vitro Diagnostic medical device as specified above conforms with the essential requirements listed in the Annex 1 of the European In Vitro Diagnostic Medical Device Directive 98/79/EC(IVDD)

This declaration of conformity is based on the European In Vitro Diagnostic Medical Device Directive98/79/EC, Annex III.

 Valid Since
 May 9th, 2012
 Representative:

 Changchun, China
 Yu Ge
 OUS TRIAL

 Dirui Industrial Cos
 Understand
 Understand

 (place and date of issue)
 (name and signature or equivalent marking of authorized person)
 (name and signature or equivalent marking of authorized person)

认证证书

标准

ISO 9001:2015

证书登记号码

01 100 1832306

证书持有者:

迪瑞医疗科技股份有限公司 统一社会信用代码: 91220101605902656F 注册地址:中华人民共和国吉林省长春市 高新技术产业开发区云河街 95 号 邮编: 130012 经营地址:同上述地址

认证范围:

体外诊断医疗器械的设计开发、生产和销售

证明完成了审核并满足了 ISO 9001:2015 标准的要求。

证书有效期从 2018-05-03 至 2021-05-02。 此证书须经过符合要求的监督审核保持有效。



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Deutsche Akkreditierungsstelle D-ZM-16031-01-00



Certificate

Standard	Standard	
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ISO 9001:2015

Certificate Registr. No.

01 100 1832306

Certificate Holder:

Dirui Industrial Co., Ltd. Unified Social Credit Code: 91220101605902656F Registration Address: 95 Yunhe Street, New & High Tech. Development Zone, Changchun City, Jilin Province 130012, P. R. China Operation Address: same as above

Scope:

Design and Development, Manufacture and Distribution of in Vitro Diagnostic Medical Test Systems

Proof has been furnished by means of an audit that the requirements of ISO 9001:2015 are met.

The certificate is valid from 2018-05-03 until 2021-05-02. It remains valid subject to satisfactory surveillance audits.



www.tuv.com



kkS Deutsche Akkreditierungsstelle D-ZM-16031-01-00





Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

Dirui Industrial Co., Ltd. 95 Yunhe Street New & High Tech. Development Zone Changchun Jilin Province 130012 China

has established and applies a quality management system for medical devices for the following scope:

Design and Development, Manufacture and Distribution of In vitro Diagnostic Medical Test Systems (see attachment for products and additional site included)

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance.

Effective Date:

2018-06-26

2020-03-01

Certificate Registration No.: SX 60127937 0001

An audit was performed. Report No.: 15047317 007

This Certificate is valid until:

Certification Body



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Date 2018-06-26

10/020 d 04.08 ®

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg Tel.: +49 221 806-1371 Fax: +49 221 806-3935 e-mail:cert-validity@de.tuv.com http://www.tuv.com/safety



Doc. 1/1, Rev. 0

TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment toCertificateRegistration No.:SX 60127937 0001Report No.:15047317 007

Organization:

Dirui Industrial Co., Ltd. 95 Yunhe Street New & High Tech. Development Zone Changchun Jilin Province 130012 China

Scope:

Products:

- Urine Test Systems (Reagents, Analyzers, Controls)
- Hematology Test Systems (Reagents, Analyzers, Controls)
- Clinical Chemistry Test Systems (Reagents, Analyzers, Controls)
- Immunochemistry Test Systems (Reagents, Analyzers, Controls)
- Vaginal Infections Test Systems (Reagents, Analyzers, Controls)

Site included:

3333 Yiju Street, New & High Tech. Development Zone, Changchun, 130103 Jilin, China

Design and Development, Manufacture and Distribution of Urine Test Analyzers, Hematology Test Analyzers, Clinical Chemistry Test Analyzers, Immunochemistry Test Analyzers, Vaginal Infections Test Analyzers

Certification Body





Date: 2018-06-26

L 10/020 d 04.08 ® TÜV, TUEV and TUV are registered trademarks. Utilisation and application requires prior approva

CE

Declaration of Conformity

Manufacturer:	Shenzhen Mindray Bio-Medical Electronics Co., Ltd.	
	Mindray Building, Keji 12th Road South, Hi-tech Industrial	
	Park, Nanshan, Shenzhen, 518057, P. R. China	
EC-Representative:	Shanghai International Holding Corp. GmbH (Europe)	
	Eiffestraße 80	
	20537 Hamburg, Germany	
Product:	Auto Hematology Analyzer	
Model:	BC-20s	
	Including reagents as following:	
	M-30D DILUENT	
	M-30CFL LYSE	
	PROBE CLEANSER	
Classification:	The device not in IVDD annex II and not for self testing/performance evaluation	

Conformity Assessment Route: IVDD Annex III(excluding Section 6)

We herewith declare that the above mentioned products meet the provisions of the Directive 98/79/EC on In Vitro Diagnostic Medical Devices. All supporting documentations are retained under the premises of the manufacturer.

Standards Applied:

List of (harmonized) standards for which documented evidence for compliance can be provided as attachment.

Start of CE-Marking: 2015-3-31 Place, Date of Issue: Shenzhen, 2015-3-31

Signature:

Name of Authorized Signatory: Mr.tan ChuanBin

Position Held in Company:

Manager, Technical Regulation

CE

Declaration of Conformity

Manufacturer:	Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
	Mindray Building, Keji 12th Road South, Hi-tech Industrial
	Park, Nanshan, Shenzhen, 518057, P. R. China
EC-Representative:	Shanghai International Holding Corp. GmbH (Europe)
	Eiffestraße 80
	20537 Hamburg, Germany
Product:	Auto Hematology Analyzer
Model:	BC-30s
	Including reagents as following:
	M-30D DILUENT M-30CFL LYSE
	PROBE CLEANSER
Classification:	The device not in IVDD annex II and not for self testing/performance evaluation

Conformity Assessment Route: IVDD Annex III(excluding Section 6)

We herewith declare that the above mentioned products meet the provisions of the Directive 98/79/EC on In Vitro Diagnostic Medical Devices. All supporting documentations are retained under the premises of the manufacturer.

Standards Applied:

List of (harmonized) standards for which documented evidence for compliance can be provided as attachment.

Start of CE-Marking: 2015-3-31

Place, Date of Issue: Shenzhen, 2015-3-31

Signature:

Name of Authorized Signatory: Mr.tan ChuanBin Position Held in Company:

Manager , Technical Regulation

Applied Standards List

Product:	Auto Hematology Analyzer
	BC-20s、BC-30s
	Including reagents as following:
	M-30D DILUENT
	M-30CFL LYSE
	PROBE CLEANSER

Applied Standards:

EN ISO 18113-1:2011	In vitro diagnostic medical devices —Information supplied by the
	manufacturer(labelling) Part 1: Terms, definitions and general requirements
ENISO 18113-2:2011	I in vitro diagnostic medical devices - Information supplied by the manufacturer
	(labelling) - Part 2: In vitro diagnostic reagents for professional use
EN ISO 18113-3:2011	In vitro diagnostic medical devices — Information supplied by the
	manufacturer(labeling) Part 3: In vitro diagnostic instruments for professional
	use
EN ISO 15223-1:2012	Medical devices Symbols to be used with medical device labels,
	labelling and information to be supplied —Part 1: General requirements
EN 13612: 2002	Performance evaluation of in vitro diagnostic medical devices
ISO 14971:2012	Medical devices – Application of risk management to medical devices
EN 61010-1:2001	Safety requirements for electrical equipment for measurement, control, and
	laboratory use Part 1: General requirement
EN 61010-2-081:2002+A1:	Safety requirements for electrical equipment for measurement, control and
2003+A1: 2003	laboratory use - Part 2-081: Particular requirements for automatic and
	semi-automatic laboratory equipment for analysis and other purposes
EN 61010-2-101: 2002	Safety requirements for electrical equipment for measurement, control, and
	laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD)
	medical equipment
IEC 61010-2-010: 2005	Safety requirements for electrical equipment for measurement, control and

laboratory use - Part 2-010: Particular requirements for laboratory equipment for the heating of materials

EN 61326-1:2006 Electrical equipment for measurement, control and laboratory use - EMC requirements - Part 1: General requirements

EN 61326-2-6:2006 Electrical equipment for measurement, control and laboratory use - EMC requirements - Part 2-6: Particular requirements - In vitro diagnostic (IVD) medical equipment

EN 62304:2006 Medical device software- Software life cycle processes

EN 62366:2008 Medical devices — Application of usability engineering to medical devices

EN 13640: 2002 Stability testing of in vitro diagnostic medical devices

EN ISO13485:2012 Medical devices - Quality management systems - Requirements for regulatory purposes







Certificate Holder:

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Mindray Building Keji 12th Road South High-Tech Industrial Park Nanshan 518057 Shenzhen PEOPLE'S REPUBLIC OF CHINA

Certification Mark:



Scope of Certificate:

Standard(s):

See Page 2 for Overall Scope Statement.

ISO 13485:2016

Regulatory Authority(ies):

Australia TGA, Brazil ANVISA, Health Canada, USA FDA, MHLW / PMDA. See attached for listing of specific regulatory requirements.

The Certification Body of TÜV SÜD America Inc. certifies that the quality management system of the manufacturer listed above has been audited against the stated criteria and found to conform to those criteria for the scope of certification listed. Validity of this certificate can be obtained by visiting the website https://www.tuev-sued.de/product-testing/certificates

TÜV SÜD America Inc. is an MDSAP Recognized Auditing Organization.

DUNS No:	65-467-1304
Effective Date:	2019-08-26
Expiry Date:	2021-10-23

Page 1 of 4 Date of Issue: 2019-11-25

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(Dawn M. Tibodeau) Manager, Certification Body MHS TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 USA • www.tuvsud.com

US-Letter / 07.17





Regulatory Reguirements:

Audit/Certification Criteria

Australia

Therapeutic Goods (Medical Devices) Regulations 2002 - Schedule 3, Part 1

Brazil

- RDC ANVISA n. 16/2013

- RDC ANVISA n. 23/2012
- RDC ANVISA n. 67/2009

Canada

- Medical Device Regulations SOR/98-282, Part 1

United States

- 21 CFR Part 803
- 21 CFR Part 806
- 21 CFR Part 807
- 21 CFR Part 820

Japan

- MHLW Ministerial Ordinance 169, Article 4 to Article 68 - PMD Act

Overall Scope Statement:

Design and Development, Production and Distribution of Medical Electronic Equipment (including Patient Monitor and Accessories (NIBP House, NIBP Cuff, Sensor Cables including SPO2 Cable and Temperature Cable, SPO2 Sensor, ECG Cables and Leadsets, Temperature Probe, Probe Cover), Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories (Vaporizer), Ventilator, Ultrasonic Diagnostic Equipment, Ultrasonic Transducer, Hematology Analyzer, Clinical Chemistry Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use, Chemiluminescence Immunossay Analyzer, Flow Cytometer, Auto Sample Processing System, Auto Slide Maker and Stainer;) Reagents for Hematology Analyzer, Reagents for Clinical Chemistry Analyzer, Chemiluminescence Immunoassay Reagents, Chemiluminescence Immunoassav Calibrators and Controls; Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, **Breathing Bag**

Page 2 of 4 Date of Issue: 2019-11-25

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(Dawn M. Tibodeau) Manager, Certification Body MHS TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 USA • www.tuvsud.com





Facility(ies):

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Mindray Building, Keji 12th Road South, High-Tech Industrial Park, Nanshan, 518057 Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. 1203 Nanhuan Avenue, Guangming District, 518106 Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Facility Scopes:

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Mindray Building, Keji 12th Road South, High-Tech Industrial Park Nanshan, 518057, Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Design and Development, Production and Distribution of Medical Electronic Equipment (including Patient Monitor and Accessories (NIBP House, NIBP Cuff, Sensor Cables including SPO2 Cable and Temperature Cable, SPO2 Sensor, ECG Cables and Leadsets, Temperature Probe, Probe Cover), Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories (Vaporizer), Ventilator, Ultrasonic Diagnostic Equipment, Ultrasonic Transducer, Hematology Analyzer, Clinical Chemistry Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use, Chemiluminescence Immunossay Analyzer, Flow Cytometer, Auto Sample Processing System, Auto Slide Maker and Stainer;) Reagents for Hematology Analyzer, Reagents for Clinical Chemistry Analyzer, Chemiluminescence Immunoassay Reagents, Chemiluminescence Immunoassav Calibrators and Controls; Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, **Breathing Bag** DUNS No: 65-467-1304

Page 3 of 4 Date of Issue: 2019-11-25

6 buon Pitrodean

(Dawn M. Tibodeau) Manager, Certification Body MHS TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 USA • www.tuvsud.com

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Shenzhen Mindray Bio-Medical Electronics Co., Ltd. 1203 Nanhuan Avenue, Guangming District, 518106, Shenzhen PEOPLE'S REPUBLIC OF CHINA

Production of Medical Electronic Equipment (including Patient Monitor and Accessories (NIBP House, NIBP Cuff, Sensor Cables including SPO2 Cable and Temperature Cable, SPO2 Sensor, ECG Cables and Leadsets, Temperature Probe, Probe Cover), Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories (Vaporizer), Ventilator, Ultrasonic Diagnostic Equipment, Ultrasonic Transducer, Hematology Analyzer, Clinical Chemistry Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use, Chemiluminescence Immunossay Analyzer, Flow Cytometer, Auto Sample Processing System, Auto Slide Maker and Stainer;) Reagents for Hematology Analyzer, Reagents for Clinical Chemistry Analyzer, Chemiluminescence Immunoassay Reagents, Chemiluminescence Immunoassav Calibrators and Controls; Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag DUNS No: 54-459-5743

Page 4 of 4 Date of Issue: 2019-11-25

6 Juin Pitrodean

(Dawn M. Tibodeau) Manager, Certification Body MHS TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 USA • www.tuvsud.com

US-Letter / 07.17







CERTIFICATE

No. QS5 044751 0140 Rev. 02

Certificate Holder:

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Mindray Building Keji 12th Road South High-Tech Industrial Park Nanshan 518057 Shenzhen PEOPLE'S REPUBLIC OF CHINA

Certification Mark:



Scope of Certificate:

See Page 2 for Overall Scope Statement.

Standard(s):

ISO 9001:2015

The Certification Body of TÜV SÜD America Inc. certifies that the company mentioned above has established and is maintaining a quality management system that meets the requirements of the listed standards.

Report No.:	SH2005501
Effective Date:	2020-08-12
Expirv Date:	2023-06-30

Page 1 of 4 Date of Issue: 2020-08-20





CERTIFICATE

No. QS5 044751 0140 Rev. 02

Overall Scope Statement

Design and Development, Production and Distribution of Medical Electronic Equipment (including Patient Monitor and Accessories, Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Temperature Probe, Flow Sensor, Ambulatory Blood Pressure Monitor, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories, Ventilator, Air Compressor, Endoscope Camera System, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use. Chemiluminescence Immunossav Analyzer. Flow Cytometer, (Auto) Sample Processing System, Auto Slide Maker and Stainer, Glycohemoglobin Analyzer, Specific Protein Analyzer), Reagents for Hematology Analyzer, Reagents for Clinical **Chemistry Analyzer, Chemiluminescence** Immunoassay Reagents, Chemiluminescence Immunoassay Calibrators and Controls, Reagents for Flow Cytometer, Reagents for Glycohemoglobin Analyzer, Calibrators and Controls for Glycohemoglobin Analyzer, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, **Disposable Breathing Circuit, Reusable Breathing** Circuit, Heat and Moisture Exchanger, Filter, **Breathing Bag**

Page 2 of 4 Date of Issue: 2020-08-20





Facility(ies):

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Mindray Building, Keji 12th Road South, High-Tech Industrial Park, Nanshan, 518057, Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Facility Scopes:

Design and Development, Production and Distribution of Medical Electronic Equipment (including Patient Monitor and Accessories, Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Temperature Probe, Flow Sensor, Ambulatory Blood Pressure Monitor, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories, Ventilator, Air Compressor, Endoscope Camera System, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use, Chemiluminescence Immunossay Analyzer, Flow Cytometer, (Auto) Sample Processing System, Auto Slide Maker and Stainer, Glycohemoglobin Analyzer, Specific Protein Analyzer), Reagents for Hematology Analyzer, Reagents for Clinical Chemistry Analyzer, Chemiluminescence Immunoassay Reagents, Chemiluminescence Immunoassay Calibrators and Controls, Reagents for Flow Cytometer, Reagents for Glycohemoglobin Analyzer, Calibrators and Controls for Glycohemoglobin Analyzer, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag

Page 3 of 4 Date of Issue: 2020-08-20





Facility(ies)

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. 1203 Nanhuan Avenue, Guangming District, 518106 Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Facility Scopes:

Design and Development, Production and Distribution of Medical Electronic Equipment (including Patient Monitor and Accessories, Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Temperature Probe, Flow Sensor, Ambulatory Blood Pressure Monitor, Defibrillator / Monitor and Accessories, Electrocardiograph, Anesthesia Machine and Accessories, Ventilator, Air Compressor, Endoscope Camera System, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for In-Vitro Diagnostic Use, Chemiluminescence Immunossay Analyzer, Flow Cytometer, (Auto) Sample Processing System, Auto Slide Maker and Stainer. Glycohemoglobin Analyzer, Specific Protein Analyzer), Reagents for Hematology Analyzer, Reagents for Clinical Chemistry Analyzer, Chemiluminescence Immunoassay Reagents, Chemiluminescence Immunoassay Calibrators and Controls, Reagents for Flow Cytometer, Reagents for Glycohemoglobin Analyzer, Calibrators and Controls for Glycohemoglobin Analyzer, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag

Page 4 of 4 Date of Issue: 2020-08-20



21.08.2016 Izmir / Turkey

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DECLARATION FOR THE ISSUANCE OF QUALITY CERTIFICATES

ITOB 10031 Sk. No:15 Menderes - Izmir / TURKEY

FACTORY | HEAD OFFICE

Sasali Merkez Mah. Doğa Dostları Sitesi 131. Sok. No:2/5 Ciğli - İzn

Tel: +90 232 376 80 81 Fax: +90 232 376 80 40

To Whom It May Concern,

According to IVD 98/79/EC directive,

FOR ANNEX II LIST A which includes HIV, Hepatitis B and Hepatitis C tests; the Notified Body must verify that the product meets the Common Technical Specification (CTS) and must release each batch of product before it is placed on the European market. The batch release often requires testing. These have EC Design Examination certificates by the notified body.

FOR ANNEX III which includes all other tests for Professional use; the manufacturer prepares a declaration of conformity in a similar way to the general devices.

For the above mentioned reason, we hereby declare that we provide CE Certificate for only the Hepatitis B, Hepatitis C and HIV tests for Professional use. For the group of other Professional tests; it is enough to present a self-Declaration of Conformity to the EU standards.

Cordially,

TURKLAB TIBBİ MALZEMELER SAN TİC A.Ş

POLSKIE CENTRUM BADAŃ I CERTYFIKACJI S.A.

POLISH CENTRE FOR TESTING AND CERTIFICATION



EC CERTIFICATE No. 1434-IVDD-56/2016

EC Design-Examination

Directive 98/79/EC on in vitro diagnostic medical devices

PCBC certifies that the design documentation relating to in vitro diagnostic medical device, List A:

HBsAg Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®

manufactured by:

TÜRKLAB Tibbi Mal. San. Tic. A.Ş. ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey

was examined by PCBC according to Annex IV p. 4 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law and comply with the essential requirements of the Directive.

This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29 Date of first certificate issue: 2008-08-29



Augeloo
Anna Wyroba
Vice President of PCBC

PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw

Application No. 45/2016 Contract No. MD-18/2016

CE 1434

Module H6

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POLSKIE CENTRUN POLSHE CENTRUN		EC CERTIFICATE No. 1434-IVDD-52/2016 EC Design-Examination	Directive 98/79/EC on in vitro diagnostic medical devices	device, List A: Anti-HCV Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®	manufactured by:	TÜRKLAB Tibbi Mal. San. Tic. A.S. ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey	was examined by PCBC according to Annex IV p. 4 Directive 98/79/BC (with subsequent amendments) transposed into the Polish law and comply with the essential requirements of the Directive.	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29	Ď	Anna Wyroba Vice President of PCBC	CE 1434 PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Application No. 43/2016 Contract No. MD-16/2016
A.		lere	DDC	REF		IDDD	ariar.	미미	CC		<u>.</u>	
POLSKIE CENTRUM BADAŃ I CERTYFIKACJI S. Polish centre for testing and certification	CERTIFICA TO AND STATE	EC CERTIFICATE No. 1434-IVDD-57/2016 Full Quality Assurance System	Directive 98/79/EC on in vitro diagnostic medical devices	TÜRKLAB Tibbi Mal. San. Tic. A.S. ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey	for the design, manufacture and final inspection of in vitro diagnostic medical devices, ^{I ist} A .	HBsAg Test HBsAg Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®	complies with the requirements of Annex IV excl. 4, 6 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law. The audit of the quality assurance system carried out by PCBC has provided evidence of the above.	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29	Date of first certificate issue: 2008-08-29	Run Wyroba Vice President of PCBC	PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Module H7
POLSKIE CENTH		EC CERTIFI	Directive 98/79	TÜR ITOB 10	for the design, manufacture	Brands: Info	complies with the re (with subsequent amendrr assurance system car	This certifia L	Dat		CE 1434	Application No. 45/2016 Contract No. MD-18/2016

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POLSKIE CENTRUM BADAŃ I CERTYFIKACJI S.A.	POLISH CENTRE FOR TESTING AND CERTIFICATION	CERTYPIC BERNARD STATE	EC CERTIFICATE No. 1434-IVDD-54/2016 ECDesign-Examination	Directive 98/79/ECon in vitro diagnostic medical devices	PCBC certifies that the design documentation relating to in vitro diagnostic medical	Anti-HBs Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®	manufactured by: TÜRKI AR Tihhi Mal San Tie, A.S.	ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey	was examined by PCBC according to Annex IV p. 4 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law and comply with the essential requirements of the Directive.	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29	Date of first certificate issue: 2008-09-29		and the second se	Anna Wyroba Vice President of PCBC	CE 1434 PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Application No. 44/2016 Contract No. MD-17/2016
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FIKACJI S.A.	RTIFICATION	CKRTYANA REALIZED TO THE CENTRE	EC CERTIFICATE No. 1434-IVDD-53/2016 Full Quality Assurance System	Directive 98/79/EC on in vitro diagnostic medical devices	PCBC certifies quality assurance system in company:	TÜRKLAB Tibbi Mal. San. Tic. A.Ş. ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey	for the design, manufacture and final inspection of in vitro diagnostic medical devices, List A:	Anti-HCV Test Brands: Info@, Toyo®, Rapidan Tester®, Labmen®	complies with the requirements of Annex IV excl. p. 4, 6 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law. The audit of the quality assurance system carried out by PCBC has provided evidence of the above.	This certificate is valid from 2016-08-29 to 2019-08-28			APP and and and and and and and and and and	Anna Wyroba Vice President of PCBC	PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Module H7
OLSKIE CENTH	FOLISH		EC CERTIF	Directive 98/79	PCBC ce	TÜR ITOB 1	for the design, manufacture	Brands: Info	complies with the rec (with subsequent amendrr assurance system ca	This certifi		Da			CE 1434	Application No. 43/2016 Contract No. MD-16/2016

		JODO	리미리	JODE	IPP	DDF				
POLSKIE CENTRUM BADAŃ I CERTYFIKACJI S.A. Polish centre for testing and certification	CERTIFIC TALE TALE TALE TALE TALE TALE TALE	ECCERTIFICATE No. 1434-IVDD-58/2016 ECDesign-Examination	Directive 98/79/EC on in vitro diagnostic medical devices PCBC certifies that the design documentation relating to in vitro diagnostic medical	device, List A: Anti - HIV 1/2 Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®	manufactured by: TÜRKI AR Tihhi Mal San Tio A S	S S S S S S S S S S S S S S S S S S S	was examined by PCBC according to Annex IV p. 4 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law and comply with the essential requirements of the Directive.	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29 Date of first certificate issue: 2008-08-29	CG 133 PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Application No. 46/2016 Contract No. MD-19/2016
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ERTYFIKACJI S.A.		EC CERTIFICATE No. 1434-IVDD-55/2016 Full Quality Assurance System	Directive 98/79/EC on in vitro diagnostic medical devices PCBC certifics quality assurance system in company:	TÜRKLAB Tibbi Mal. San. Tic. A.Ş. ITOB 10031 Sokak No: 15 Tekeli Menderes Izmir, Turkey	for the design, manufacture and final inspection of in vitro diagnostic medical devices, List A:	Anti-HBs Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen®	complies with the requirements of Annex IV excl. p. 4, 6 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law. The audit of the quality assurance system carried out by PCBC has provided evidence of the above.	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29 Date of first certificate issue: 2008-08-29	PCBC Notified Body PCBC Notified Body 23A, Klobucka Str., PL-02-699 Warsaw	Module H7
POLSKIE CENTRUM B/ POLSKIE CENTRUM B/	THE CENTRE	EC CERTIFICATE Full Quality	Directive 98/79/EC on in vi PCBC certifies quality as	TÜRKLAB Tibł ITOB 10031 Sokak ¹ Izmi	for the design, manufacture and final inspe 1	Anti- Brands: Info®, Toyo®, R	complies with the requirements of An (with subsequent amendments) transposed assurance system carried out by PCF	This certificate is valid fron Date of certificat Date of first certific	CE 1434 23A, Ki	Application No. 44/2016 Contract No. MD-17/2016

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POLSKIE CENTRUM BADAŃ I CERTYFIKACJI S.A. POLSH CENTRE FOR TESTING AND CERTFICATION	EC CERTIFICATE No. 1434-IVDD-51/2016 EC Design-Examination Directive 98/79/EC on in vitro diagnostic medical devices	PCBC certifies that the design documentation relating to in vitro diagnostic medical device for self-testing: hCG Pregnancy Test Brands: Rapidan Nova@, Rapidan Optima@, Info@, Toyo@, Rapidan Tester@, Rapidan Compact@, Labmen@ manufactured hv	TÜRKLAB ITOB 10031 So was examined by PCBC aco (with subsequent amendments) tt	This certificate is valid from 2016-08-29 to 2019-08-28 Date of certificate issue: 2016-08-29 Date of first certificate issue: 2008-08-29		Application No. 42/2016 Contract No. MD-15/2016 Module A1
POLSKIE CENTRUM BADAN I CERTYFIKACJI SA. Polski centre for testing and certification	EC CERTIFICATE No. 1434-IVDD-59/2016 Full Quality Assurance System Directive 98/79/EC on in vitro diagnostic medical devices	e system in company: L San. Tic. A.S. Tekeli Menderes rikey fin vitro diagnostic medical devices,	List A: Anti - HIV 1/2 Test Anti - HIV 1/2 Test Brands: Info®, Toyo®, Rapidan Tester®, Labmen® complies with the requirements of Annex IV excl. p. 4, 6 Directive 98/79/EC (with subsequent amendments) transposed into the Polish law. The audit of the quality assurance system carried out by PCBC has provided evidence of the above.		C € 143 23A, Klobucka Str., PL-02-699 Warsaw	Application No. 46/2016 Contract No. MD-19/2016



CERTIFICATE

No. J - 2670/2/2018

This is to certify that:

TÜRKLAB TIBBI MALZ. SAN. VE TIC. A.Ş. Sasalı Merkez Mh. Doğa Dostları Sitesi 131 Sk. No: 2/5 35621 Çiğli, İzmir, Turkey

Factory: ITOB 10031 Sk. No: 15 Menderes / İzmir - Turkey

is in conformance with

EN ISO 9001:2015

in the following scope of activities:

design, development, manufacturing, final control and distribution of in vitro diagnostic medical devices intended for self-testing and professional use, ECG electrodes and antibiotic susceptibility discs

The audit carried out by the Polish Centre for Testing and Certification has afforded evidence of the above. This Certificate shall remain valid provided that above standard are respected by the Organization.

This certificate is valid:

from 24.08.2018 to 21.12.2020









Polish Centre for Testing and Certification 23A Kłobucka Street, 02-699 Warsaw Poland, phone +48 22 46 45 200, e-mail:pcbc@pcbc.gov.pl



Certificate No. J - 2670/2/2018 Issued under the Contract No. 2897/JM/3/2017 Date of certification decision: 24.08.2018 Bears the PCBC hologram. Warsaw, 24.08.2018



CERTIFICATE

No. M - 56/2/2018

This is to certify that:

TÜRKLAB TIBBI MALZ. SAN. VE TIC. A.Ş. Sasalı Merkez Mh. Doğa Dostları Sitesi 131 Sk. No: 2/5 35621 Çiğli, İzmir, Turkey Factory: ITOB 10031 Sk. No: 15 Menderes / İzmir - Turkey

is in conformance with

EN ISO 13485:2016

in the following scope of activities:

design, development, manufacturing, final control and distribution of in vitro diagnostic medical devices intended for self-testing and professional use, ECG electrodes and antibiotic susceptibility discs

The audit carried out by the Polish Centre for Testing and Certification has afforded evidence of the above. This Certificate shall remain valid provided that above standard are respected by the Organization.

This certificate is valid:

from 24.08.2018 to 21.12.2020









Polish Centre for Testing and Certification 23A Kłobucka Street, 02-699 Warsaw Poland, phone +48 22 46 45 200, e-mail:pcbc@pcbc.gov.pl



Certificate No. M - 56/2/2018 Issued under the Contract No. 2897/JM/3/2017 Date of certification decision: 24.08.2018 Bears the PCBC hologram. Warsaw, 24.08.2018