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ORDIN DE PLATA NR.: 1049 TIP.DOC. 1 :
DATA EMITERII:13 decembrie 2021 :
=====:
PLATITI: 2600-00 LEI: Doua Mii Sase Sute lei 00 bani :
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:
=====:
PLATITOR: (R) "BIOSISTEM CONTUL DE PLATI/CODUL IBAN :
MLD" S.R.L. MD95ML000000002251429243 :
CODUL FISCAL :1010600028048 / :
:
:
=====:
PRESTATORUL PLATITOR CODUL BANCII:
BC"Moldindconbank"S.A. suc."Invest" Chisinau :MOLDMD2X329:
=====:
BENEFICIAR (R) IMSP SR CAH CONTUL DE PLATI/CODUL IBAN :
UL MD39VI0000000022514095MDL :
CODUL FISCAL :1009603003860 / :
:
:
=====:
PRESTATORUL BENEFICIAR CODUL BANCII:
B.C."VICTORIABANK"S.A. :VICBMD2X :
=====:
DESTINATIA PLATII: Pentru garantia pentru: TIPUL TRANSFERULUI :
oferta la procedura de achizitie public: NORMAL/URGENT :N:
a nr. ocds-b3wdp1-MD-1637735939304 din 1: :
4.12.2021 :
:
:
L.S. :
=====:
CODUL TRANZACTIEI:001: :
DATA PRIMIRII:13/12/2021 : SEMNATURILE :
DATA EXECUTARII: : EMITENTULUI :
-----:
CONducATOR:Web Poiata Vitalie :
MIIGYwYJKoZIhvcNAQcCoIIGVDCcBlACAQExCzAJBgUrDgMCGGUAMAsGCSqGSIB3:
DQEHAAcCBGwwggRoMIIDUKADAgECAhNHAACjbilrgFksQ0G4AAAAAKNuMA0GCSqG:
SIB3DQEBcWUAMCIXIDAeBgNVBAMTF0NFU1QxLUNBLU1vbGRpbmRjb25iYW5rMB4X:
DTIxMDEyODExMzgWNVVoXDTI0MDEyODExNDgwNVowgZ8xCzAJBgNVBAYTAk1EMRAw:
YDVQQIEwdNb2xkb3ZhmREwDwYDVQQHEwhDaGlzaW5hdTEWMBQGA1UEChMNQmlv :
:
(semnatura electronica) :
CONTABIL-SEF:Web Nasedchin Alexandr :
MIIGZwYJKoZIhvcNAQcCoIIGWDCCBlQCAQExCzAJBgUrDgMCGGUAMAsGCSqGSIB3:
DQEHAAcCBHAWggRsMIIDVKADAgECAhNHAACjcahRKqbJeg8QAAAAAKNxMA0GCSqG:
SIB3DQEBcWUAMCIXIDAeBgNVBAMTF0NFU1QxLUNBLU1vbGRpbmRjb25iYW5rMB4X:
DTIxMDEyODExMzkxOFoXDTI0MDEyODExNDkxOFowgAMxMzAJBgNVBAYTAk1EMRAw:
YDVQQIEwdNb2xkb3ZhmREwDwYDVQQHEwhDaGlzaW5hdTEWMBQGA1UEChMNQmlv :
:
L.S. (semnatura electronica) :
CONducATOR: :
(semnatura manuala) :
CONTABIL-SEF: :
(semnatura manuala) :
SEMnATURA PRESTATORUL L.S. :
:
MOTIVUL REFUZULUI : L.S. :
-----:

**CERTIFICAT**  
**privind lipsa sau existența restanțelor față de bugetul public național**

Nr.  
№ **A2120445**

din  
от **26.11.2021**

**1. Destinația / Назначение**

Pentru participarea la proceduri de achiziții publice

**2. Date despre contribuabil / Информация о налогоплательщике**

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

**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 11.12.2021**

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



**Sef DDF Rîșcani**  
**Șef DCAF mun. Chișinău**

L.S./М.П. **Natalia Cebanova**

Executor: \_\_\_\_\_  
Numele și prenumele/Фамилия и имя

\_\_\_\_\_   
Semnătura/Подпись

**Viorica CĂUȘ**

\_\_\_\_\_   
Numele și prenumele/Фамилия и имя

Este extras din Sistemul Informațional al SFS SIA „Contul curent al contribuabilului”// 26.11.2021 ora 15:30:27  
cu aplicarea prevederilor pct. 82-83 Ordin IFPS nr.400 din 14.03.2014 (Monitorul Oficial 72-77/399, 28.03.2014)

NOTA (3,52)



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

Republica Moldova, MD-2068  
mun. Chişinău, bd. Moscovei, 14/1  
Tel. : (373-22) 43-44-81, 43-46-24  
Fax : (373-22) 43-44-22  
cod: MOLDM2X329

Data 14. IAN. 2016  
Nr. 03/2 - 19/23

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: MOLDM2X329

Filiala „Invest” BC „Moldindconbank” SA confirmă existența contului curent  
in moneda nationala al **“BIOSISTEM MLD” S.R.L. (c/f 1010600028048)**, cu  
**IBAN MD95ML000000002251429243.**

Codul băncii MOLDM2X329.

Director

Nina Turcan

Director financiar



Nina Balmuş

Ex. Diana Brinza  
Tel. 43-45-96

REPUBLICA



MOLDOVA

# CERTIFICAT DE ÎNREGISTRARE

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

*Numărul de identificare de stat - codul fiscal*  
**1010600028048**

*Data înregistrării*

**12.08.2010**

*Data eliberării*

**12.08.2010**

**Svirepova Ludmila, registrator**

*Funcția, numele, prenumele persoanei  
care a eliberat certificatul*

*L. Svirepova*  
semnătura

MD 0101250





**I.P. "AGENȚIA SERVICII PUBLICE"**

Departamentul înregistrare și licențiere a unităților de drept

**EXTRAS**

**din Registrul de stat al persoanelor juridice**

nr. 8506 din 28.04.2021

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.**

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,**

Asociați:

- 1. POIATA VITALIE 33,40 %**
- 2. NASEDCHIN ALEXANDR 33,30 %**
- 3. KOJEVNIKOV DMITRII 33,30 %.**

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: 28.04.2021.

Specialist coordonator  
tel. 022-207-840



**Lazari Aliona**



EB 0358735

## **Lista fondatorilor Biosistem-mld SRL**

<b>Nr.</b>	<b>Nume, Prenume</b>	<b>IDNP</b>
<b>1.</b>	<b>Vitalie Poiata</b>	<b>0983103892591</b>
<b>2.</b>	<b>Alexandr Nasedchin</b>	<b>2002001070747</b>
<b>3.</b>	<b>Dmitrii Kojevnikov</b>	<b>0972305012362</b>

Declaration of Conformity V 1.0

# 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-5150  
Including reagents as following:  
**M-52D DILUENT**  
**M-52DIFF LYSE**  
**M-52LH 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:** 2013-9-26

**Place, Date of Issue:** Shenzhen, 2013-9-26

**Signature:** \_\_\_\_\_

**Name of Authorized Signatory:** Mr.tan ChuanBin  
**Position Held in Company:** Manager ,Technical Regulation

Declaration of Conformity V 1.0

## Applied Standards List

**Product:** Auto Hematology Analyzer  
BC-5150、BC-5000  
Including reagents as following:  
**M-52D DILUENT**  
**M-52DIFF LYSE**  
**M-52LH LYSE**  
**PROBE CLEANSER**

### Applied Standards:

EN ISO 18113-1:2009	In vitro diagnostic medical devices —Information supplied by the manufacturer(labelling) Part 1: Terms, definitions and general requirements
EN ISO 18113-2:2009	In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 2: In vitro diagnostic reagents for professional use
EN ISO 18113-3:2009	In vitro diagnostic medical devices — Information supplied by the manufacturer (labelling) 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: 2003+A1: 2003	Safety requirements for electrical equipment for measurement, control and 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



## Declaration of Conformity V 1.0

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 62304:2008	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

TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD TÜV SÜD  
ZERTIFIKAT ♦ CERTIFICATE ♦ 認證證書 ♦ CERTIFICADO ♦ CERTIFICAT



# Certificate

No. Q5 044751 0164 Rev. 02

**Holder of Certificate:** **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:**

**Design and development, production and distribution of Active medical devices (intended) for monitoring, diagnosis, anesthesia, breathing and intensive care; In-vitro diagnostic instruments; Non-active accessories for breathing therapy and anesthesia; In-vitro diagnostic reagents and kits (intended) for hematology, clinical chemistry, immunology and cell analysis (For detail information see following pages)**

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). See also notes overleaf.

**Report No.:** SH2005501

**Valid from:** 2020-09-01  
**Valid until:** 2023-08-31

**Date,** 2020-07-24

Christoph Dicks  
Head of Certification/Notified Body

# Certificate

No. Q5 044751 0164 Rev. 02

**Applied Standard(s):** EN ISO 13485:2016  
Medical devices - Quality management systems -  
Requirements for regulatory purposes  
(ISO 13485:2016)  
DIN EN ISO 13485:2016

**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



# Certificate

No. Q5 044751 0164 Rev. 02

For the product(s)/product category (ies):

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 invitro diagnostic use, Chemiluminescence Immunoassay Analyzer, Flow Cytometer, (Auto) Sample Processing System, Auto Slide Maker&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.



America

# 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

**Expiry Date:** 2023-06-30

Page 1 of 4

**Date of Issue:** 2020-08-20

Tina Israel  
Manager, US Certification Body,  
Medical and Health Services



America

# 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 Immunoassay 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

Tina Israel  
Manager, US Certification Body,  
Medical and Health Services



America

# CERTIFICATE

No. QS5 044751 0140 Rev. 02

**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 Immunoassay 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

Tina Israel  
Manager, US Certification Body,  
Medical and Health Services



America

# CERTIFICATE

No. QS5 044751 0140 Rev. 02

**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 Immunoassay 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

Tina Israel  
Manager, US Certification Body,  
Medical and Health Services





# H-100

Urine Analyzer



**DIRUI**

# H-100

Urine Analyzer

## Product Characteristics:

- Adopting the advanced high luminosity cold light source with 4 -wavelength, which improves the sensitivity, accuracy, specificity, and reduces the interference from ambient light
- Adopting automatic waste handling system, which avoids cross-contamination between samples
- Automatically rectifies the test results influenced by non-specificity, pH, specific gravity, and color
- Built-in thermal printer with high speed and low noise; External stylus printer
- Connectable with DIRUI urine sediment analyzer
- Users can set an abnormal value flag by themselves
- International, regular and symbol system units display for option



## Technical Specification:

- Test items: urobilinogen, bilirubin, ketone, Creatinine, Micro-albumin, blood, protein, nitrite, leukocytes, glucose, specific gravity, pH and VC
- Test wavelength: 525nm, 572nm, 610nm, 660nm
- Test principle: Photoelectric colorimetry
- Suitable strips: DIRUI H8, H10, H11, H11-MA, H13-Cr, Urinalysis strips
- Test throughput: 120 strips/h or 60 strips/h optional
- Data storage: 1000 patient results
- Computer interface: RS-232 port; parallel printer interface
- Display: 240×64 dot-matrix LCD
- Language: Chinese, English, Russian, Polish, Italian, Spanish, Portuguese, Turkish, German, French
- Power supply: 100~240VAC, 50Hz/60Hz
- Power: 40VA
- Dimensions: 376mm×316mm×170mm
- Weight: 3.6kg
- Printer: Built-in thermal printer



Certified to  
ISO 9001:2008 and ISO 13485:2003

## DIRUI INDUSTRIAL CO., LTD.

95 Yunhe Street, New & High Tech. Development Zone  
Changchun, Jilin 130012 P. R. China

Tel: +86 431 85100409 Fax: +86 431 85172581

E-mail: dirui@dirui.com.cn Http://www.dirui.com.cn

· Specifications subject to change without notice.

20130510

# DIRUI

**C E R T I F I C A T E**  
of Conformity



Registration No.: AK 50205311 0001

Report No.: 16800459 001

Holder: Dirui Industrial Co., Ltd.  
95 Yunhe Street  
New & High Tech. Development Zone  
Changchun, Jilin 130012  
P.R. China

Product: Analysis Equipment  
(Urine Analyzer)

Identification: Type Designation: H-50 H-100 H-300 H-500  
Serial No.: Engineering Sample  
Remark: Refer to test report 16800459 001 for details.

Tested acc. to: EN 61326-1:2006  
EN 61326-2-6:2006

The certificate of conformity refers to the above mentioned product. This is to certify that the specimen is in conformity with the assessment requirement mentioned above. This certificate does not imply assessment of the production of the product and does not permit the use of a TÜV Rheinland mark of conformity.

Date 24.06.2011



Certification Body

A handwritten signature in blue ink, appearing to read "Sun Lixun".

Sun Lixun

**TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg**



# Declaration of Conformity



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 Representative:** Emergo Europe

Molenstraat 15 2513 BH The Hague  
The Netherlands

**Medical Device :** Product Name: Reagent strips for Urinalysis

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
- DIRUI A10    DIRUI H10    DIRUI E10    DIRUI M10    DIRUI H10-800
- DIRUI H11    DIRUI H11-MA    DIRUI H11-800
- DIRUI H11-800MA      DIRUI H12-800MA
- DIRUI H13-Cr      DIRUI H14-Ca
- DIRUI H13-Cr (H-800)      DIRUI H14-Ca (H-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 Directive 98/79/EC, Annex III.**

Valid Since  
May 9<sup>th</sup>, 2012  
Changchun, China

Representative:  
Yu Ge  
Dirui Industrial Co., Ltd.  
  


(place and date of issue)

(name and signature or equivalent marking of authorized person)

# 证书附件

标准 **ISO 9001:2015**  
证书登记号码 **01 100 1832306**

号码	场地	认证范围
/01	(分证书) 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	体外诊断医疗器械的设计开发、生产和销售
/02	(分证书) 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：中华人民共和国吉林 省长春市高新技术产业开发区 宜居路 3333 号 邮编：130103	体外诊断医疗器械的设计开发、生产和销售

2021-04-19

  
TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

页 1 / 1

# 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

所包括场地已列于证书附件上

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。

有效期: 证书有效期从 2021-05-03 至 2024-05-02。  
此证书须经过符合要求的监督审核保持有效。  
初次发证始于 2018 年  
本证书信息可在国家认证认可监督管理委员会官方网站上查询  
<http://www.cnca.gov.cn>

2021-04-19

TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# 认证证书

标准 **ISO 9001:2015**

证书登记号码 **01 100 1832306/01**

主证持有者: **迪瑞医疗科技股份有限公司**  
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

场地: **(分证书) 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。

有效期: 证书连同主证书 01 100 1832306 一起有效期从 2021-05-03 至 2024-05-02。

此证书须经过符合要求的监督审核保持有效。  
本证书信息可在国家认证认可监督管理委员会官方网站上查询  
<http://www.cnca.gov.cn>

2021-04-19

TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# 认证证书

标准 **ISO 9001:2015**

证书登记号码 **01 100 1832306/02**

主证持有者: **迪瑞医疗科技股份有限公司**  
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

场地: **(分证书) 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: 中华人民共和国吉林省长春市  
高新技术产业开发区宜居路 3333 号  
邮编: 130103

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。

有效期: 证书连同主证书 01 100 1832306 一起有效期从 2021-05-03 至 2024-05-02。

此证书须经过符合要求的监督审核保持有效。  
本证书信息可在国家认证认可监督管理委员会官方网站上查询  
<http://www.cnca.gov.cn>

2021-04-19

TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln



# Annex to certificate

Standard **ISO 9001:2015**

Certificate Registr. No. **01 100 1832306**

No.	Location	Scope
/01	c/o Dirui Industrial Co., Ltd. Unified Social Credit Code: 91220101605902656F Registration Address: 95 Yunhe Street, New & High Tech. Development Zone, Changchun, 130012 Jilin, P. R. China Operation Address: same as above	Design and Development, Manufacture and Sales of In Vitro Diagnostic Medical Test Systems
/02	c/o Dirui Industrial Co., Ltd. Unified Social Credit Code: 91220101605902656F Registration Address: 95 Yunhe Street, New & High Tech. Development Zone, Changchun, 130012 Jilin, P. R. China Operation Address: 3333 Yiju Street, New & High Tech. Development Zone, Changchun, 130103 Jilin, P. R. China	Design and Development, Manufacture and Sales of In Vitro Diagnostic Medical Test Systems

2021-04-19

  
TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# Certificate

Standard **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, 130012 Jilin, P. R. China  
Operation Address: same as above

including the locations according to annex

Scope: Design and Development, Manufacture and Sales 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.

Validity: The certificate is valid from 2021-05-03 until 2024-05-02.  
It remains valid subject to satisfactory surveillance audits.  
First certification 2018  
This certificate information can be searched on CNCA official  
website <http://www.cnca.gov.cn>

2021-04-19



TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# Certificate

Standard **ISO 9001:2015**

Certificate Registr. No. **01 100 1832306/01**

Organization: **Dirui Industrial Co., Ltd.**  
95 Yunhe Street, New & High Tech. Development Zone,  
Changchun, 130012 Jilin, P.R. China

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

Scope: Design and Development, Manufacture and Sales 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.

Validity: The certificate is valid in conjunction with the main certificate 01  
100 1832306 from 2021-05-03 until 2024-05-02.  
It remains valid subject to satisfactory surveillance audits.  
This certificate information can be searched on CNCA official  
website <http://www.cnca.gov.cn>

2021-04-19



TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# Certificate

Standard **ISO 9001:2015**

Certificate Registr. No. **01 100 1832306/02**

Organization: **Dirui Industrial Co., Ltd.**  
95 Yunhe Street, New & High Tech. Development Zone,  
Changchun, 130012 Jilin, P. R. China

Site: **c/o Dirui Industrial Co., Ltd.**  
Unified Social Credit Code: 91220101605902656F  
Registration Address: 95 Yunhe Street,  
New & High Tech. Development Zone,  
Changchun, 130012 Jilin, P. R. China  
Operation Address: 3333 Yiju Street,  
New & High Tech. Development Zone,  
Changchun, 130103 Jilin, P. R. China

Scope: Design and Development, Manufacture and Sales 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.

Validity: The certificate is valid in conjunction with the main certificate 01  
100 1832306 from 2021-05-03 until 2024-05-02.  
It remains valid subject to satisfactory surveillance audits.  
This certificate information can be searched on CNCA official  
website <http://www.cnca.gov.cn>

2021-04-19

TÜV Rheinland Cert GmbH  
Am Grauen Stein · 51105 Köln

# Certificate

## Quality Management System EN ISO 13485:2016

Registration No.: SX 2101045-1

Organization: Dirui Industrial Co., Ltd.  
95 Yunhe Street, New & High Tech. Development Zone, Changchun,  
130012 Jilin, P.R. China

Scope: Design and Development, Manufacture and Distribution of In-vitro Diagnostic Analyzers and In-Vitro Diagnostic Test Kits used in the Detection of Blood Analyte, Cancer, Cardiac Markers, Coagulation Blood, Endocrine Disorders, Immune Status, Protein Metabolism, Sexually Transmitted Diseases, Infectious Disease, Therapeutic Drug Monitoring, Disease Status, Renal Function Assessment, Liver Function Assessment, Pancreas Function Assessment, Blood Fat Test, Physiological Markers, Nucleic Acid Extraction Reagent and Specimen Receptacle for Clinical Laboratory Use.

The Certification Body of TÜV Rheinland LGA Products GmbH certifies that the organization has established and applies a quality management system for medical devices. Proof has been furnished that the requirements specified in the abovementioned standard are fulfilled. The quality management system is subject to yearly surveillance.

Report No.: 190131562 110  
Effective date: 2021-04-30  
Expiry date: 2023-03-01  
Issue date: 2021-05-08



Jing Zhang  
TÜV Rheinland LGA Products GmbH  
Tillystraße 2 · 90431 Nürnberg · Germany

# Certificate

**Quality Management System  
EN ISO 13485:2016**

Registration No.: SX 2101045-1

Organization: Dirui Industrial Co., Ltd.  
95 Yunhe Street, New & High Tech. Development Zone, Changchun,  
130012 Jilin, P.R. China

The scope of certification also covers the following:

No.	Facility	Scope
/01	c/o Dirui Industrial Co., Ltd. 95 Yunhe Street, New & High Tech. Development Zone, Changchun, 130012 Jilin, P.R. China	Design and Development, Manufacture of In-Vitro Diagnostic Test Kits used in the Detection of Blood Analyte, Cancer, Cardiac Markers, Coagulation Blood, Endocrine Disorders, Immune Status, Protein Metabolism, Sexually Transmitted Diseases, Infectious Disease, Therapeutic Drug Monitoring, Disease Status, Renal Function Assessment, Liver Function Assessment, Pancreas Function Assessment, Blood Fat Test, Physiological Markers, Nucleic Acid Extraction Reagent and Specimen Receptacle for Clinical Laboratory Use.

Report No.: 190131562 110  
Effective date: 2021-04-30  
Expiry date: 2023-03-01  
Issue date: 2021-05-08



Jing Zhang  
TÜV Rheinland LGA Products GmbH  
Tillystraße 2 · 90431 Nürnberg · Germany

# Certificate

**Quality Management System  
EN ISO 13485:2016**

Registration No.: SX 2101045-1

Organization: Dirui Industrial Co., Ltd.  
95 Yunhe Street, New & High Tech. Development Zone, Changchun,  
130012 Jilin, P.R. China

The scope of certification also covers the following:

/02 c/o Dirui Industrial Co., Ltd.  
3333 Yiju Street, New & High Tech.  
Development Zone, Changchun,  
130103 Jilin, P.R. China

Design and Development, Manufacture and Distribution of In-vitro Diagnostic Analyzers used in the Detection of Blood Analyte, Cancer, Cardiac Markers, Coagulation Blood, Endocrine Disorders, Immune Status, Protein Metabolism, Sexually Transmitted Diseases, Infectious Disease, Therapeutic Drug Monitoring, Disease Status, Renal Function Assessment, Liver Function Assessment, Pancreas Function Assessment, Blood Fat Test, Physiological Markers for Clinical Laboratory Use.

Report No.: 190131562 110  
Effective date: 2021-04-30  
Expiry date: 2023-03-01  
Issue date: 2021-05-08



Jing Zhang  
TÜV Rheinland LGA Products GmbH  
Tillystraße 2 · 90431 Nürnberg · Germany

# EC DECLARATION OF CONFORMITY

*In vitro* Diagnostic Medical Devices in accordance with Directive 98/79/EC

**Manufacturer:** Türklab Tıbbi Mal. San. ve Tic. A.Ş.  
**Headquarters / Manufacturing Side:** İTOB 10017 Sokak No: 2 Tekeli - Menderes / İzmir - Turkey  
**Product:** Troponin I Test  
**Brand:** Rapidan® Tester, Toyo®, Info®, Labmen®  
**Classification:** Professional Use IVD, 98/79/EC  
**Conformity Assessment Route:** Annex III

We, herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for *In-Vitro* Diagnostic Medical Devices. All supporting documentation is retained under the premises of the manufacturer.

**Standards Applied:** EN ISO 13485:2016  
EN ISO 14971:2012  
EN ISO 15223:2016  
EN ISO 18113-1:2011  
EN ISO 18113-2:2011  
EN ISO 23640:2015  
EN 13612:2002

**Revision No:** 4

**Place, Date of Issue:** İzmir, 08.03.2019

**Signature** Dr. Şahin Yağlıdere, Md  
General Manager

**TÜRKLAB**  
TIBBİ MALZ. SAN. VE TİC. A.Ş.  
MERKEZ: İTOB OSB MAH. 10031 SK. NO: 15 MENDERES / İZMİR  
FABRİKA: İTOB OSB MAH. 10017 SK. NO: 2 MENDERES / İZMİR  
TEL: 0 232 376 80 81 - FAX: 0 232 376 80 40  
MENDERES V.D. 879 009 6209







# CERTIFICATE

**No M - 56/4/2020**

This is to certify that:

**TÜRKLAB Tibbi Mal. San. Tic. A.Ş**  
**ITOB 10017 Sokak No: 2,**  
**Tekeli - Menderes İzmir / Turkey**

and

**Location**

**listed in Annex to the certificate**

is in conformance with

**EN ISO 13485:2016**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
rapid tests intended for self-testing and for professional use,  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents) and ECG electrodes**

The audit carried out by the Polish Centre of 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 **22.12.2020** to **21.12.2023**

Issued under the Contract No. 2897/JM/4/2020

Date of certification decision: 14.10.2020

Certificate bears a qualified signature.

Warsaw, 15.10.2020



AC 019  
QMS



Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
09:00:16 +02'00'



# ANNEX TO THE CERTIFICATE

VALID ONLY IN CONNECTION WITH THE CERTIFICATE

**No M - 56/4/2020**

This is to certify that the following Location:

**Factory 2: ITOB 10031 Sokak No: 15,  
Tekeli - Menderes İzmir / Turkey**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents),  
professional use IVD tests and ECG electrodes**

**meets the requirements of the standard listed on the certificate**

Issued under the Contract No. 2897/JM/4/2020  
Date of certification decision: 14.10.2020  
Certificate bears a qualified signature.  
Warsaw, 15.10.2020



AC 019  
QMS



Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
09:02:27 +02'00'



# CERTIFICATE

**No J - 2670/4/2020**

This is to certify that:

**TÜRKLAB Tibbi Mal. San. Tic. A.Ş.**  
**ITOB 10017 Sokak No: 2,**  
**Tekeli - Menderes İzmir / Turkey**

and

**Location**

**listed in Annex to the certificate**

is in conformance with

**EN ISO 9001:2015**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
rapid tests intended for self-testing and for professional use,  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents) and ECG electrodes**

The audit carried out by the Polish Centre of 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 **22.12.2020 to 21.12.2023**

Issued under the Contract No. 2897/JM/4/2020

Date of certification decision: 14.10.2020

Certificate bears a qualified signature.

Warsaw, 15.10.2020



AC 019  
QMS



Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
08:47:33 +02'00'



# ANNEX TO THE CERTIFICATE

VALID ONLY IN CONNECTION WITH THE CERTIFICATE

**No J - 2670/4/2020**

This is to certify that the following Location:

**Factory 2: ITOB 10031 Sokak No: 15,  
Tekeli - Menderes İzmir / Turkey**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents),  
professional use IVD tests and ECG electrodes**

**meets the requirements of the standard listed on the certificate**

Issued under the Contract No. 2897/JM/4/2020  
Date of certification decision: 14.10.2020  
Certificate bears a qualified signature.  
Warsaw, 15.10.2020



AC 019  
QMS



Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
08:48:40 +02'00'

SOĞUK DAMGA VARDIR



T.C.  
TORBALI 6. NOTERLİĞİ  
Telf: 0232 664 70 07 Fax: 0232 664 70 17



№ 09971

## SERTİFİKA

No. M – 56/4/2020

İşbu sertifika ile;

**TÜRKLAB Tıbbi Mal. San. Tic. A.Ş.**  
**ITOB 10017 Sokak No:2, Tekeli-Menderes**  
**İzmir, Türkiye**

ve sertifika ekinde listelenmiş

### Lokasyon

Aşağıdaki faaliyetler kapsamında

**EN ISO 13485:2016**

ile uyumludur:

**invitro tıbbi cihazların tasarımı, geliştirilmesi, üretimi, son kontrolü ve dağıtımı: kendi kendine test ve profesyonel kullanım için tasarlanmış hızlı testler, kan gruplaması için reaktifler ve reaktif ürünleri (jel kartları ve kırmızı kan hücreleri reaktifleri) ve EKG elektrotları**

Polonya Test ve Sertifikasyon Merkezi tarafından yürütülen denetim, yukarıdaki kanıtları sağlamıştır. Bu Sertifika, Kuruluş tarafından yukarıdaki standarda uyulması kaydıyla geçerliliğini koruyacaktır.

**Bu sertifikanın geçerlilik tarihi: 22.12.2020'den 21.12.2023'e kadar**

Sözleşme Çerçevesinde Düzenleme No.2897/JM/4/2020

Sertifika kararının tarihi: 14.10.2020

Sertifika, yetkili imzayı taşımaktadır.

Varşova, 15.10.2020

Anna <<Elektronik İmza>>  
Malgorzata  
Wyroba  
Yönetim Kurulu Üyesi

POLONYA TEST VE SERTİFİKASYON MERKEZİ 02-844 Varşova, 469 Pulawska Street, Tel: +48 22 46 45 200, e-posta: [pcbc@pcbc.gov.pl](mailto:pcbc@pcbc.gov.pl)

İşbu belge İngilizce aslından Türkçe'ye tarafımdan aslına uygun olarak tercüme edilmiştir.

I hereby certify that this document has been translated from its English into Turkish truly and correctly by me. 03.12.2020

SWORN TRANSLATOR / YEMİNLİ TERCÜMAN  
ERKAN ALTUNER

10 3 Aralık 2020





# CERTIFICATE

No M - 56/4/2020

This is to certify that:

**TÜRKLAB Tibbi Mal. San. Tic. A.Ş**  
**ITOB 10017 Sokak No: 2,**  
**Tekeli - Menderes İzmir / Turkey**

and

**Location**

**listed in Annex to the certificate**

is in conformance with

**EN ISO 13485:2016**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
rapid tests intended for self-testing and for professional use,  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents) and ECG electrodes**

The audit carried out by the Polish Centre of 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 22.12.2020 to 21.12.2023

Issued under the Contract No. 2897/JM/4/2020  
Date of certification decision: 14.10.2020  
Certificate bears a qualified signature.  
Warsaw, 15.10.2020

T.C.  
TORBALI 6. NOTERİ  
Selma ZİYREK

Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
09:00:16 +02'00'

PARTNER OF  
**IONet**



AC 019  
QMS





№ 09971

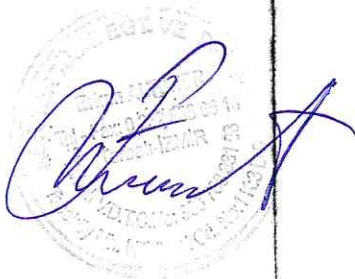
**SERTİFİKA EKİ****SADECE SERTİFİKA İLE BAĞLANTILI OLARAK GEÇERLİDİR**  
**No. M – 56/4/2020**

İşbu sertifika, aşağıda yer alan faaliyetler kapsamında Lokasyonun tasdiki için hazırlanmıştır:

**Fabrika 2: ITOB 10031 Sokak No: 15,**  
**Tekeli-Menderes İzmir, Türkiye****invitro tıbbi cihazların tasarımı, geliştirilmesi, üretimi, son kontrolü ve dağıtımı: kan gruplaması için reaktifler ve reaktif ürünleri (jel kartları ve kırmızı kan hücreleri reaktifleri), profesyonel kullanım IVD testleri ve EKG elektrotları****Sertifikada listelenen standardın gereksinimlerini karşılar.**Sözleşme Çerçevesinde Düzenleme No.2897/JM/4/2020  
Sertifika kararının tarihi: 14.10.2020  
Sertifika, yetkili imzayı taşımaktadır.  
Varşova, 15.10.2020Anna <<Elektronik İmza>>  
Malgorzata  
Wyroba  
Yönetim Kurulu ÜyesiPOLONYA TEST VE SERTİFİKASYON MERKEZİ 02-844 Varşova, 469 Pulawska Street, Tel: +48 22 46 45 200, e-posta: [pcbc@pcbc.gov.pl](mailto:pcbc@pcbc.gov.pl)

İşbu belge İngilizce aslından Türkçe'ye tarafımdan aslına uygun olarak tercüme edilmiştir.

I hereby certify that this document has been translated from its English into Turkish truly and correctly by me. 03.12.2020

**SWORN TRANSLATOR / YEMİNLİ TERCÜMAN**  
ERKAN ALTUNER

03 Aralık 2020





## ANNEX TO THE CERTIFICATE

VALID ONLY IN CONNECTION WITH THE CERTIFICATE

**No M - 56/4/2020**

This is to certify that the following Location:

№ 09971

**Factory 2: ITOB 10031 Sokak No: 15,  
Tekeli - Menderes İzmir / Turkey**

in the following scope of activities:

**design, development, manufacturing, final control  
and distribution of in vitro medical devices:  
reagents and reagent products for blood grouping  
(gel cards and red blood cells reagents),  
professional use IVD tests and ECG electrodes**

meets the requirements of the standard listed on the certificate

Issued under the Contract No. 2897/JM/4/2020  
Date of certification decision: 14.10.2020  
Certificate bears a qualified signature.  
Warsaw, 15.10.2020



AC 019  
QMS



Anna  
Małgorzata  
Wyroba  
Member of the Board

Elektronicznie  
podpisany przez Anna  
Małgorzata Wyroba  
Data: 2020.10.16  
09:02:27 +02'00'

T.C.  
TORBALI 6. NOTER  
Seim ZIYREK

