

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

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: MOLDMD2X329

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

1 Balney

Codul băncii MOLDMD2X329.

Director

Director financia

Nina Turcan

Nina Balmuş

Ex. Diana Brinza Tel. 43-45-96



CERTIFICAT DE ÎNRECISTRARE

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 S. Sizes

MD 0101250





I.P. "AGENŢIA 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



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. | Alexandr Nasedchin | 2002001070747 |
| 3. | Dmitrii Kojevnikov | 0972305012362 |

CERTIFICAT privind lipsa sau existența restanțelor față de bugetul public național

| Nr. _№ A2201651 din _{от} 02.02.2022 | | | | |
|---|--|--|--|--|
| 1. Destinația / Назначение | | | | |
| Pentru participare 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) Соdul - Denumirea localității Код - Наименование населенного пункта | | | | |
| Albisoara nr.16 bl.1 of.7 0150-S | EC.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 17.02.2022 | | | | |
| 5. Autentificarea Serviciului Fiscal de Stat / Подтверждение Государственной налоговой службы | | | | |
| Sef DDF Riscani Semnätura/Подпись L.S/MIF Executor: Svettana Slonovscaia Numelest prenumele/Фамилия и имя 1022)823102 | Numele și prenumele/Фамилия и имя | | | |

Este extras din Sistemul Informațional al SFS SIA "Contul curent al contribuabilului"// 02.02.2022 ora 8:34:43 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)



EC DECLARATION OF CONFORMITY

Apacor Limited declares that the devices listed below conform to the relevant provisions of the EC Council Directive In Vitro Diagnostic Devices Directive 98/79/EC dated 27 October 1998. This compliance has been properly documented using checklist created from Annex III excluding point 6 of the Directive, linked to all supporting Technical Documentation.

Apacor Limited has a Quality Management System in place, which complies with ISO 13485 (Certificate Number GB18/873854) regulations and agrees to develop, implement and maintain the Quality Management System to ensure continued adequacy and efficacy.

| PRODUCT DESCRIPTION | PRODUCT CODE | EDMS CODE | CATEGORY |
|----------------------|---|-----------|--------------------|
| MIDI PARASEP | 145000, 145300, 145400, 145500, 145501, 145650, 145750, 249200 | 15051090 | Other Parasitology |
| MIDI PARASEP SF | 149900, 149910, 149920, 149931, 149932, 149650, 149750, 249300 | 15051090 | Other Parasitology |
| MINI PARASEP | 146000, 146200, 146300, 146400, 146500, 146501, 146650, 146750, 248200 | 15051090 | Other Parasitology |
| MINI PARASEP SF | 148800, 148900, 148910, 148920, 148931, 148932, 148935, 148980, 148650, 148750, 248930,108000,180880, 108810,108900,108910,108920,108931, 108932,108935 | 15051090 | Other Parasitology |
| MAXI PARASEP | 147001 | 15051090 | Other Parasitology |
| 30ML TRANSPORT VIALS | 148998, 249400, 249420 | 15051090 | Other Parasitology |
| CLEAN VIAL | 149970 | 15051090 | Other Parasitology |

This Declaration of Conformity is signed below, certifying these requirements have been met.

Janet MacKenzie General Manager

15 September 2018



EC Certificate

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

Registration No.: HL 60131743 0001

Report No.: 10042449 010

Manufacturer: Bioptik Technology, Inc.

No. 188, Jhonghua South Road

Gongguan Village Jhunan Township Miaoli County, 35057

Taiwan

Products: In-vitro diagnostic Medical Devices for self-testing

(see attachment for products included)

Replaces Approval, Registration No.: HL 60088590 0001

Expiry Date: 2023-09-17

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

Effective Date: 2018-10-19

Date: 2018-10-19

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg
TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 98/79/EC
concerning in vitro diagnostic medical devices with the identification number 0197.



TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Doc. 1/1, Rev. 0

Attachment to Certificate

Registration No.: Report No.:

HL 60131743 0001 10042449 010

Manufacturer:

Bioptik Technology, Inc.

No. 188, Jhonghua South Road

Gongguan Village Jhunan Township Miaoli County, 35057

Taiwan

Products:

- Blood Glucose Monitoring Systems
- Blood Cholesterol Monitoring Systems
- Hemoglobin Monitoring Systems
- Blood Triglyceride Monitoring System
- Blood Glucose/Uric Acid Monitoring Systems
- Blood Glucose/Cholesterol Monitoring Systems
- Blood Glucose/Hemoglobin Monitoring System
- Blood Glucose/Cholesterol/Uric Acid Monitoring Systems
- Blood Glucose/Cholesterol/Hemoglobin Monitoring Systems
- Blood Glucose/Cholesterol/Triglyceride Monitoring Systems
- Blood Glucose/Cholesterol/Uric Acid/Hemoglobin Monitoring System
- Blood Glucose/Cholesterol/Uric Acid/Hemoglobin/ Triglyceride Monitoring Systems
- Blood Pressure/Glucose/Cholesterol Monitoring Systems (assessment limited to Glucose/Cholesterol Monitoring)

Date: 2018-10-19





Certificate

The Certification Body of TÜV Rheinland LGA Products GmbH

hereby certifies that the organization

Bioptik Technology, Inc. No. 188, Jhonghua South Road Gongguan Village Jhunan Township Miaoli County, 35057 Taiwan

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

Design and development, manufacture and distribution of Medical devices (see attachment for products 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:

2018-10-19

Certificate Registration No.:

SX 60131746 0001

An audit was performed. Report No.: 50145079 001

This Certificate is valid until:

2021-09-17

Certification Body



Date 2018-10-19



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



TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Doc. 1/1, Rev. 0

Attachment to Certificate

Registration No.: SX 60131746 0001 Report No.: 50145079 001

Organization:

Bioptik Technology, Inc.

No. 188, Jhonghua South Road

Gongguan Village Jhunan Township Miaoli County, 35057

Taiwan

Scope:

Products:

- In vitro diagnostic medical devices used in blood analytes and blood glucose monitoring including meter, test strips and control solutions for self-testing, near patient/point of care.
- Blood Pressure/Glucose/Cholesterol Monitoring System (assessment limited to Blood Pressure Monitoring)

Certification Body



Date: 2018-10-19



Certificate

Standard

ISO 9001:2015

Certificate Registr. No.

01 100 6696

Certificate Holder:

BIOSYSTEMS 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





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

Certification Body



Date 2020-01-08

Torrheinland Land Control of the Con

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



TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Doc. 1/1, Rev. 0

Attachment to Certificate

Registration No.: SX 60145545 0001 Report No.: 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

DAKKS

Deutsche
Akkreditierungsstelle
D-ZM-14169-01-02

Date: 2020-01-08

Certification Body

D. Swiatko





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

Dr. Antonio Elduque Managing director BioSystems S.A.





CLINICAL CHEMISTRY - BIOCHEMISTRY:

a-Amylase-Direct a-Amylase-EPS

a-Amylase-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)

Calcium-Arsenazo

Bilirubin (total)

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-Keratin Antibodies (AKA)
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 (ASMA)

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/RK/RS (AA-RL/RK/RS)
Glomerular Basement Membrane
Antibodies (GBMA)



AUTOIMMUNITY - ELISA:

ANA Screening
Anti-Annexin V IgG/IgM (ANX)
Anti-b2-Glycoprotein 1 IgG/IgM
(b2GP1)
Anti-Cardiolipin Antibodies (ACA-IgG/IgM)
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

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: ITOB 10017 Sokak No: 2 Tekeli - Menderes / Izmir - Turkey

Fecal Occult Blood (FOB) Test Product:

Rapidan® Tester, Toyo®, Info®, Labmen® Brand:

Professional Use IVD, 98/79/EC Classification:

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: 5

Place, Date of Issue: Izmir, 08.03.2019

Signature Dr. Şahin Yağlıdere, Md

General Manager

MERKEZ: İTOB OSB MAH. 10081 51/4 MO:15 MENDERES / IZMİR FABRİKA: İTOB OSB MAH 10011/21/4. NO:2 MENDERES / IZMİR TEL: 0 232 376 80 81 FAX: 0 232 376 80 40 MENDERES V.D/379 009 6209



No J-2670/4/2020

This is to certify that:

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

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 affored 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







Member of the Board



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 | zmir / 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







Member of the Board

TERCÜME

T.C. ORBALI 6. NOTERLIĞI 0/232 554 70 07 Fax: 0/232 604 70 17

N209971

SERTIFIKA

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 SERTIFIKASYON MERKEZİ 02-844 Varşova, 469 Pulawska Street, Tel: +48 22 46 45 200, e-posta: pcbc@pcbc.gov.pl

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

I herebycertifythatthisdocument has beentranslatedfromits English intoTurkishtrulyandcorrectlyby me.03.12.2020

SWORN TRANSLATOR / YEMINLI TERCUMAN

ERKAN ALTUNER

10 3 Wally 5050

TORBALI 6 NOTERI-Selme ZIVREK



No M-56/4/2020

This is to certify that:

TÜRKLAB Tibbi Mal. San. Tic. A.Ş ITOB 10017 Sokak No: 2, Tekeli - Menderes | zmir / 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 cellsreagents) and ECG electrodes

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

This certificate is valid:

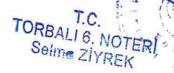
from 22.12.2020 to 21.12.2023

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

MO 9 9 1





Anna / Małgorzata Elektronicznie podpisany przez Anna Małgorzata Wyroba Data: 2020.10.16

Wyroba 09:00:16 ±02'00'
Member of the Board









Nº09971

SERTIFIKA EKI

SADECE SERTIFIKA İLE BAĞLANTILI OLARAK GEÇERLİDİR No. M – 56/4/2020

İşbu sertifika, aşağıda yer alan faaliyetler kapsamındaLokasyonun tasdiki için hazırlanmıştır:

Fabrika 2:ITOB 10031 Sokak No: 15, Tekeli-Menderesİzmir, 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, 15.10.2020

Anna <<Elektronik İmza>>

Malgorzata Wyroba

Yönetim Kurulu Üyesi

POLONYA TEST VE SERTIFIKASYON MERKEZİ 02-844 Varşova, 469 Pulawska Street, Tel: +48 22 46 45 200, e-posta: pcbc@pcbc.gov.pl

İşbu belge İngilizce aslından Türkçe'ye tarafımdan aslına uygun olarak tercüme edilmiştir. I herebycertifythatthisdocument has beentranslatedfromits English intoTurkishtrulyandcorrectlyby me.03.12.2020

SWORN TRANSLATOR / YEMINLI TERCÜMAN

ERKAN ALTUNER

0 3 Malik 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:

Factory 2: ITOB 10031 Sokak No: 15, Tekeli - Menderes | zmir / 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







Anna

TORBALI 6. NOT

Wyroba

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

Member of the Board

Page 1 of 1



No M - 56/4/2020

This is to certify that:

TÜRKLAB Tibbi Mal. San. Tic. A.Ş ITOB 10017 Sokak No: 2, Tekeli - Menderes | zmir / 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 cellsreagents) and ECG electrodes

The audit carried out by the Polish Centre of Testing and Certification has affored 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







Member of the Board



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 | zmir / 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







Member of the Board

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-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:

Signature:

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

Name of Authorized Signatory: Mr.tan ChuanBin

Position Held in Company: Manager ,Technical Regulation

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

Including reagents as following:

BC-30s

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:

Signature:

Model:

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

Mr.tan ChuanBin

Start of CE-Marking: 2015-3-31

Name of Authorized Signatory:

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

Position Held in Company: Manager ,Technical Regulation

Declaration of Conformity V 1.0

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

| Declaration of Conformity V 1.0 | | | | |
|---------------------------------|---|--|--|--|
| | 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 | | | |







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





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

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





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 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

Tina Israel

Manager, US Certification Body, Medical and Health Services





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 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

Tina Israel

Manager, US Certification Body, Medical and Health Services







Product Service

Certificate

No. Q5 044751 0164 Rev. 02

Holder of Certificate: Shenzhen Mindray Bio-Medical

Electronics Co., Ltd.

Mindray Building Keii 12th Road South High-Tech Industrial Park

Nanshan

518057 Shenzhen

PEOPLE'S REPUBLIC OF CHINA

Certification Mark:



Design and development, **Scope of Certificate:**

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.

SH2005501 Report No.:

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

2020-07-24

Christoph Dicks

Head of Certification/Notified Body

Date,





Certificate

No. Q5 044751 0164 Rev. 02

EN ISO 13485:2016 Applied Standard(s):

Medical devices - Quality management systems -

Requirements for regulatory purposes

(ISO 13485:2016) DIN EN ISO 13485:2016

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Facility(ies):

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.