

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







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)
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-Keratin Antibodies (AKA)
Anti-Mitochondrial Antibodies (AMA)
Anti-Noutrophil 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/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-Histones Antibodies (HIST)

Anti-Insulin Antibodies (INS)

Anti-Jo1 Antibodies

Anti-M2 Antibodies (M2)

Anti-Gliadin Antibodies (AGA-IgG/IgA)

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:

BIOSYSTEMS S.A.

Costa Brava, 30 08030 Barcelona

Spain

(including the locations according to annex)

Scope:

Design, development, manufacture, distribution, installation and

servicing of:

- Instruments and reagents for clinical diagnostic.

- Instruments and reagents for agro-alimentary analysis. Distribution and servicing of instruments and reagents 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 2017-12-13 until 2019-12-18.

First certification 1996

2017-12-14

TÜV Rheinland Cert GmbH Am Grauen Stein · 51105 Köln









Annex to certificate

Standard

ISO 9001:2015

Certificate Registr. No.

01 100 6696

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N	\mathbf{a}	

Location

Scope

/01

BIOSYSTEMS, S.A.

PI. Can Tapioles naus 7-12-13 08110 Montcada i Reixac

Spain

Labelling and assembling of reagents.

Warehousing and shipment

of:

-Instruments and Reagents

for clinical diagnostic.

-Instruments and Reagents for agro-alimentary analysis. -Instruments and Reagents

for veterinary diagnosis.

2017-12-14

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:

2017-11-28

Certificate Registration No.:

SX 60124804 0001

An audit was performed. Report No.: 28300434 002

This Certificate is valid until:

2019-12-12

Certification Body



Date 2017-11-28



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 to Certificate

Registration No.: Report No.:

SX 60124804 0001 28300434 002

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 (Barcelona)

Spain

Scope:

Labelling and Assembling of reagents and Warehousing and Shipment of instruments and

reagents for clinical diagnostic

Certification Body



Date: 2017-11-28





Declaration of Conformity

This is to state that Technical Documentation (CL001, rev. 2.0) for product(s)

Coaguometer

(Model:CA-01, CA-02)

(IVD products other than those covered by Annex II, IVD for self-testing and devices for Performance evaluation according to manufacturer's declaration)

Manufactured by

CLINDIAG SYSTEMS CO., LTD

No.29 Zhiyuan Road, Jurong Economic Development Zone, Zhenjiang, Jiangsu Province, China

Has been assessed as meeting the Essential Requirements and relevant provisions of EC Directive 98/79/EEC for in Vitro Diagnostic





REGISTRATION NO. 04716Q10011R0S

CERTIFICATE OF QUALITY MANAGEMENT SYSTEM

This is to certify that the quality management system of

Clindiag Systems Co., Ltd.

Registered Address: No.29 Zhiyuan Road, Jurong Economic Development Zone, Jiangsu Province, P.R.China Postcode: 212400

Manufacturing Address: No.29 Zhiyuan Road, Jurong Economic Development Zone, Jiangsu Province, P.R.China

Has been assessed and conformed to the following standard(s)
GB/T 19001-2016 idt ISO 9001: 2015

The certificate is valid for the following scope:

The Design, Development, Production and Service of
Semi-Automatic Biochemistry Analyzer, Coagulometer Analyzer,
Semi-Automatic Electrolyte Analyzer, Semi-Automatic Microplate
Reader, Microplate Washer, Automatic Urine Analyzer And Urine
Test Strips, Fully Automatic Biochemistry Analyzer, Fully
Automatic Hematology Analyzer, Vitro Diagnostic Reagent

(for export Only)

Date of issue: January 15, 2019
Date of expiry: January 14, 2021

Director: Throng hen

BEIJING HUA GUANG CERTIFICATION OF MEDICAL DEVICES CO., LTD.



MANAGEMENT SYSTEM CNAS C047 - Q





REGISTRATION NO. 04716Q10000011

CERTIFICATE OF QUALITY MANAGEMENT SYSTEM FOR MEDICAL DEVICES

This is to certify that the quality management system of

Clindiag Systems Co., Ltd.

Registered Address: No.29 Zhiyuan Road, Jurong Economic Development

Zone, Jiangsu Province, P.R.China Postcode: 212400

Manufacturing Address: No.29 Zhiyuan Road, Jurong Economic

Development Zone, Jiangsu Province, P.R.China

Has been assessed and conformed to the following standard(s) YY/T 0287-2017 idt ISO 13485: 2016

The certificate is valid for the following scope:

The Design, Development, Production and Service of Semi-Automatic Biochemistry Analyzer, Coagulometer Analyzer, Semi-Automatic Electrolyte Analyzer, Semi-Automatic Microplate Reader, Microplate Washer, Automatic Urine Analyzer And Urine Test Strips, Fully Automatic Biochemistry Analyzer, Fully Automatic Hematology Analyzer, Vitro Diagnostic Reagent.

(for export Only)

Date of issue: January 15, 2019

Date of expiry: January 14, 2021

Director: Juganglann

BEIJING HUA GUANG CERTIFICATION OF MEDICAL DEVICES CO., LTD.



www.imq.it

CERTIFICATO N. CERTIFICATE N. 9190.CRC3



IQNet, the association of the world's first class certification bodies, is the largest provider of management System Certification in the world. IONet is composed of more than 30 hodies and counts over 150 subsidiaries all over the globe.

SI CERTIFICA CHE IL SISTEMA QUALITA' DI WE HEREBY CERTIFY THAT THE QUALITY SYSTEM OPERATED BY

CERACARTA SPA

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

UNITA' OPERATIVE / OPERATIVE UNITS

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

E' CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

ISO 9001:2008

PER LE SEGUENTI ATTIVITA' / FOR THE FOLLOWING ACTIVITIES

Produzione e stampa di carte speciali e diagrammate per registrazione ad uso industriale, ferroviario, medicale e biglietteria anche conto terzi. Produzione e stampa di etichette e biglietti anche a lettura/scrittura in radio frequenza (RFID). Sviluppo e produzione di creme, gel sterile e non sterile per applicazioni elettrodiagnostiche e ad ultrasuoni, anche conto terzi. Commercializzazione ed immissione in commercio di accessori per applicazioni elettrodiagnostiche, ad ultrasuoni e per strumenti elettromedicali. Sviluppo e produzione di elettrodi per ECG. Gestione della produzione ed immissione in commercio di elettrodi per ECG. Immissione in commercio di piastre per elettrobisturi e defibrillatori. Commercializzazione di video stampanti, carte fotografiche per video stampanti, stampanti e relativi materiali di consumo ed accessori Manufacture and print of special recording chart papers for industrial, railway, medical use and ticketing also on behalf of third parties. Manufacture and print of labels and tickets also radio frequency reading/writing (RFID). Development and manufacture of creams, gels sterile and not sterile for electromedical and ultrasound procedures also on behalf of third parties. Trade and placing on the market of accessories for electromedical and ultrasound diagnostic devices and for electromedical equipment. Development and manufacture of electrods for ECG. Production management and placing on the market of electrods for ECG. Placing on the market of electrosurgical plates and defibrillation pads. Trade of videoprinters, photographic papers for videoprinters, printers and related consumable and accessories

Ulteriori informazioni riguardanti l'applicabilità dei requisiti ISO 9001:2008 possono essere ottenute consultando l'organizzazione Further clarifications regarding the applicability of ISO 9001:2008 requirements may be obtained by consulting the organization

> IL PRESENTE CERTIFICATO E' SOGGETTO AL RISPETTO DEL REGOLAMENTO PER LA CERTIFICAZIONE DEI SISTEMI DI GESTIONE THE USE AND THE VALIDITY OF THE CERTIFICATE SHALL SATISFY THE REQUIREMENTS OF THE RULES FOR CERTIFICATION OF MANAGEMENT SYSTEMS

DATE:

PRIMA CERTIFICAZIONE FIRST CERTIFICATION

IAF: 07, 09, 19, 29

EMISSIONE CORRENTE **CURRENT ISSUE**

SCADENZA EXPIRY

2002-11-26

2017-10-13

2020-10-07

L'Organizzazione dovrà ottenere la certificazione secondo la norma ISO 9001:2015 entro il 2018/09/14; in caso contrario, il presente certificato cesserà la propria validità in tale data The Organization shall obtain the certification according to ISO 9001:2015 within 2018/09/14; otherwise the validity of this certificate will expire

> IMQ S.p.A. - VIA QUINTILIANO, 43 - 20138 MILANO ITALY Management Systems Division - Flavio Ornago

Data di scadenza del precedente ciclo di certificazione: 2017-10-07

Data di conclusione dell'audit di rinnovo: 2017-10-11

Data della decisione di rinnovo: 2017-10-13





www.cisq.com

Organismo di Certificazione Federato CISQ

CISO è la Federazione Italiana di Organismi di

CREDIA



THE INTERNATIONAL CERTIFICATION NETWORK

CERTIFICATE

CISQ/IMQ as an IQNet Partner hereby states that the organization

CERACARTA SPA

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

for the following scope:

Manufacture and print of special recording chart papers for industrial, railway, medical use and ticketing also on behalf of third parties. Manufacture and print of labels and tickets also radio frequency reading/writing (RFID). Development and manufacture of creams, gels sterile and not sterile for electromedical and ultrasound procedures also on behalf of third parties. Trade and placing on the market of accessories for electromedical and ultrasound diagnostic devices and for electromedical equipment. Development and manufacture of electrods for ECG. Production management and placing on the market of electrods for ECG. Placing on the market of electrosurgical plates and defibrillation pads. Trade of videoprinters, photographic papers for videoprinters, printers and related consumable and accessories

Further clarifications regarding the applicability of ISO 9001:2008 requirements may be obtained by consulting the organization

has implemented and maintains a

Quality Management System

which fulfills the requirements of the following standard

ISO 9001:2008

Issued on: 2017 - 10 - 13

First issued on: 2002 - 11 - 26

for the validity date, please refer to the original certificate* issued by IMO

Registration Number: IT – 112265



Alex Stoichitoiu President of IQNET



Ing. Claudio Provetti President of CISO

IQNet Partners**:

AENOR Spain AFNOR Certification France APCER Portugal CCC Cyprus CISQ Italy COC China COM China COS Czech Republic Cro Cert Croatia DQS Holding GmbH Germany FCAV Brazil FONDONORMA Venezuela ICONTEC Colombia Inspecta Certification Finland INTECO Costa Rica IRAM Argentina JQA Japan KFQ Korea MIRTEC Greece MSZT Hungary Nemko AS Norway NSAI Ireland PCBC Poland Quality Austria Austria RR Russia SIGE México SII Israel SIQ Slovenia SIRIM QAS International Malaysia SQS Switzerland SRAC Romania TEST St Petersburg Russia TSE Turkey Vincotte Belgium YUQS Serbia IONet is represented in the USA by: AFNOR Certification, CISQ, DQS Holding GmbH and NSAI Inc.

* This attestation is directly linked to the IQNet Partner's original certificate and shall not be used as a stand-alone document ** The list of IONet partners is valid at the time of issue of this certificate. Updated information is available under www.iqnet-certification.com



CERTIFICATO N. CERTIFICATE N. 9124.CRC4



IQNet, the association of the world's first class certification bodies, is the largest provider of management System Certification in the world. IQNet is composed of more than 30 bodies and counts over 150 subsidiaries all over the globe.

SI CERTIFICA CHE IL SISTEMA QUALITA' DI WE HEREBY CERTIFY THAT THE QUALITY SYSTEM OPERATED BY

CERACARTA SPA

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

UNITA' OPERATIVE / OPERATIVE UNITS

VIA SECONDO CASADEI 14 Z.I. VILLA SELVA - 47122 FORLI' (FC)

E' CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

EN ISO 13485:2012

PER LE SEGUENTI ATTIVITA' / FOR THE FOLLOWING ACTIVITIES

Produzione e stampa di carte speciali e diagrammate per registrazione ad uso medicale anche conto terzi. Produzione e stampa di etichette ad uso medicale. Sviluppo e produzione di creme, gel sterile e non sterile per applicazioni elettrodiagnostiche e ad ultrasuoni, anche conto terzi. Commercializzazione ed immissione in commercio di accessori per applicazioni elettrodiagnostiche, ad ultrasuoni e per strumenti elettromedicali. Sviluppo e produzione di elettrodi per ECG. Gestione della produzione ed immissione in commercio di elettrodi per ECG. Immissione in commercio di piastre per elettrobisturi e defibrillatori. Commercializzazione di video stampanti, carte fotografiche per video stampanti, stampanti e relativi materiali di consumo ed accessori per uso medicale

Manufacture and print of special recording chart papers for medical use also on behalf of third parties. Manufacture and print of labels for medical use. Development and manufacture of creams, gels sterile and not sterile for electromedical and ultrasound procedures also on behalf of third parties. Trade and placing on the market of accessories for electromedical and ultrasound diagnostic devices and for electromedical equipment. Development and manufacture of electrods for ECG. Production management and placing on the market of electrods for ECG. Placing on the market of electrosurgical plates and defibrillation pads. Trade of videoprinters, photographic papers for videoprinters, printers and related consumable and accessories for medical use

Ulteriori informazioni riguardanti l'applicabilità dei requisiti EN ISO 13485:2012 possono essere ottenute consultando l'organizzazione Further clarifications regarding the applicability of EN ISO 13485:2012 requirements may be obtained by consulting the organization

IL PRESENTE CERTIFICATO E' SOGGETTO AL RISPETTO DEL REGOLAMENTO PER LA CERTIFICAZIONE DEI SISTEMI DI GESTIONE

THE USE AND THE VALIDITY OF THE CERTIFICATE SHALL SATISFY THE REQUIREMENTS OF THE RULES FOR CERTIFICATION OF MANAGEMENT SYSTEMS

DATE:

PRIMA CERTIFICAZIONE FIRST CERTIFICATION

1999-07-20

EMISSIONE CORRENTE **CURRENT ISSUE**

2017-10-13

SCADENZA **EXPIRY**

2020-10-07

L'Organizzazione dovrà ottenere la certificazione secondo la norma ISO 13485:2016 entro il 2019/02/28; in caso contrario, il presente certificato cesserà la propria validità in tale data The Organization shall obtain the certification according to ISO 13485:2016 within 2019/02/28; otherwise the validity of this certificate will expire

> IMQ S.p.A. - VIA QUINTILIANO, 43 - 20138 MILANO ITALY Management Systems Division - Flavio Ornago

Data di scadenza del precedente ciclo di certificazione: 2017-10-07 Data di conclusione dell'audit di rinnovo: 2017-10-11 Data della decisione di rinnovo: 2017-10-13

CISQ è la Federazione Italiana di Organismi di Certificazione dei sistemi di gestione aziendale.

CISQ is the Italian Federation of management system Certification Bodies.



www.cisq.com



Declaration of Conformity CE

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: Model:

Auto Hematology Analyzer

BC-3600

Including reagents as following

M-30D DILUENT

M-30CFL LYSE

M-30R RINSE

PROBE CLEANSER

Classification:

he device not in IVDD annex II and not for self

testing/performance evaluation

Conformity Assessment Route: WDD 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: 2011-01-14

Place, Date of Issue: Shenzhen, 2011-01-14

Signature:

Name of Authorized Signatory:

Mr. Yang Long

Position Held in Company:

Management Representative

DAKKS CRT2 / 10.13



CERTIFICATE

No. Q5 17 03 44751 089

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.

SH1705528 Report No.:

2017-09-01 Valid from: Valid until: 2020-08-31

Date. 2017-06-28 Stefan Preiß









CERTIFICATE No. Q5 17 03 44751 089

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. Bldg 9-13, Baiwangxin High-Tech Industrial Park, Baimang, Xili Town, Nanshan, 518108 Shenzhen,

PEOPLE'S REPUBLIC OF CHINA

Shenzhen Mindray Biomedical Electronics Co., Ltd. 1203 Nanhuan Avenue, Guangming District, 518106

Shenzhen, PEOPLE'S REPUBLIC OF CHINA







Attachment for Certificate No. Q5 17 03 44751 089

Dated: 2017-06-28

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, Wearable ECG Recorder.

Anesthesia Machine and Accessories, Ventilator,

Infusion Pump, Syringe Pump, Enteral Feeding Pump, Infusion Supervision System,

Ultrasonic Diagnostic Equipment and Accessories,

Digital Radiography System, Radiography System, Magnetic Resonance Imaging System

Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader,

Microplate Washer for invitro diagnostic use, Chemiluminescence Immunossay 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

Munich, CRT, 2017-06-28

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Stefan Preiß

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CERTIFICATE

No. QS5 17 07 44751 097

Certificate Holder:

Shenzhen Mindrav 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 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, Wearable ECG Recorder, Anesthesia Machine and Accessories, Ventilator, Infusion Pump, Syringe Pump, Enteral Feeding Pump, Infusion Supervision System, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Magnetic Resonance Imaging System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for Invitro Diagnostic Use, Chemiluminescence Immunossay 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 Immunoassav 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,

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.:

M2606

Effective Date:

2017-07-01

Expiry Date:

2020-06-30

Earl Buckmiller

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Director, Quality Systems & MS Cert. Body

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TÜV SÜD America Inc. 10 Centennial Drive Peabody, MA 01960 USA





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CERTIFICATE

No. QS5 17 07 44751 097

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, Vital Signs Monitor, Center Monitoring System, Telemetry Monitoring System, Pulse Oximeter, Temperature Probe, Flow Sensor, Ambulatory Blood pressure Monitor, Defibrillator/Monitor and Accessories, Electrocardiograph, Wearable ECG Recorder, Anesthesia Machine and Accessories, Ventilator, Infusion Pump, Syringe Pump, Enteral Feeding Pump, Infusion Supervision System, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Magnetic Resonance Imaging System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for Invitro Diagnostic Use, Chemiluminescence Immunossay 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 Immunoassav 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

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. Bldg 9-13, Baiwangxin High-Tech Industrial Park Baimang, Xili Town Nanshan, 518108 Shenzhen PEOPLE'S REPUBLIC OF CHINA

Design and Development, Manufacturing of 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, Wearable ECG Recorder, Anesthesia Machine, Ventilator, Ultrasonic Diagnostic Equipment and Accessories, Digital Radiography System, Radiography System, Magnetic Resonance Imaging System. Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag

Effective Date:

2017-07-01

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2020-06-30

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TÜV SÜD America Inc. 10 Centennial Drive Peabody, MA 01960 USA







CERTIFICATE

No. QS5 17 07 44751 097

Shenzhen Mindray Bio-Medical Electronics Co., Ltd. 1203 Nanhuan Avenue Guangming District 518016 Shenzhen PEOPLE'S REPUBLIC OF CHINA

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, Wearable ECG Recorder, Anesthesia Machine and Accessories, Ventilator, Infusion Pump, Syringe Pump, Enteral Feeding Pump, Infusion Supervision System, Ultrasonic Diagnostic Equipment and Accessories (Ultrasonic Transducer), Digital Radiography System, Radiography System, Magnetic Resonance Imaging System, Hematology Analyzer, Clinical Chemistry Analyzer, Urine Analyzer, Microplate Reader, Microplate Washer for Invitro Diagnostic Use, Chemiluminescence Immunossay 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 Immunoassav 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

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