

"Echipamed-Plus" SRL str. Valea Trandafirilor, 24B, of. 2-7 MD-2001, Chisinau, Moldova +373 22 234-349

Date: 03.12.2021

#### **LETTER OF AUTHORIZATION**

To whom it may concern,

We, Shenzhen Mindray Bio-Medical Electronics Co., Ltd., ("Mindray") manufacturer of CL-900i, CL-1000i, CL-1200i, BS-230, BS-240pro, BS-430, BS-800, corresponding reagents and consumables ("Products"), hereby certify that we authorize "Echipamed-Plus" SRL, with business office at str. Valea Trandafirilor, 24B, of. 2-7, MD-2001, Chisinau, Republic of Moldova ("You") as the exclusive distributor and local representative for sales and service of the Products in Republic of Moldova ("Territory").

As the manufacturer, Mindray guarantees the Products against defects in materials and workmanship, and provide services based on the standard terms and conditions of Mindray's warranty policy.

This authorization of distribution rights is valid from the date of issuance to **December 31**, **2022**. Mindray reserves the right to terminate the authorization upon fifteen (15) days written notice without any compensation to You.

Neither this Letter of Authorization nor any further extension, will impose any obligation or grant any rights regarding further distribution of the Products, nor allow any party to seek compensation for goodwill developed during the term of Letter of Authorization or any further extension.

Best regards.

General Manager of Sales and Warketing Division, CIS

Shenzhen Mindray Bio Medical Electronics Co., Ltd.

SHENZHEN MINDRAY BIO-MEDICAL ELECTRONICS CO., LTD.

Duan Lian

Mindray Building, Kejr 12th Road South, High-tech Industrial Park, Nanshan, Shenzhen 518057, P.R. China Tel: +86 755 81888998

Fax: +86 755 26582680 Website: www.mindray.com









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

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Date of Issue: 2020-08-20

Tina Israel

Manager, US Certification Body, Medical and Health Services

TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 USA •

TUV®

www.tuvsud





#### 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 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** 

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Date of Issue: 2020-08-20

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

TÜV SÜD America Inc. • 10 Centennial Drive Ste 207 • Peabody, MA 01960 US









Product Service

### 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

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TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich •



### 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, Keii 13th Road South, High-Tech I

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



TÜV®





#### 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.



# 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: Chemiluminescence Immunoassay Analyzer

Model: CL-1000i

Consumables: Reaction cuvettes.

waste container

ptional Module:

Built-in sample bar code reader

Optional Module: Built-in sample bar code reader

Built-in reagent bar code reader

Classification: The device not in IVDD annex II and not for self

testing/performance evaluation

Conformity Assessment Route: IVDD Annex III (not includes 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.

Mr. Tan Chuanbin

Start of CE-Marking: 2015-09-30

Name of Authorized Signatory:

Place, Date of Issue: Shenzhen, 2015-09-30

Position Held in Company: Manager of Technical Regulation

# 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:

Chemiluminescence Immunoassay Analyzer

Model:

CL-1200i

Consumables:

Reaction cuvettes.

waste container

Optional Module:

Built-in sample bar code reader

Built-in reagent bar code reader

Hand-held bar code reader

Classification:

The device not in IVDD annex II and not for self

testing/performance evaluation

Conformity Assessment Route: IVDD Annex III (not includes Section 6)

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

Standards Applied:

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

Start of CE-Marking: 2015-09-30

Place, Date of Issue:

Shenzhen, 2015-09-30

Signature:

Name of Authorized Signatory:

Mr. Tan Chuanbin

Position Held in Company:

Manager of Technical Regulation

#### **Applied Standards List**

**Product:** 

Chemiluminescence Immunoassay Analyzer

CL-1000i /CL-1200i

**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

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

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 Safety requirements for electrical equipment for measurement, control and

+A1: 2003 laboratory use - Part 2-081: Particular requirements for automatic and

semi-automatic laboratory equipment for analysis and other purposes

EN 61010-2-101: 2002 Safety requirements for electrical equipment for measurement, control, and

laboratory use - Part 2-101: Particular requirements for in vitro diagnostic

(IVD) medical equipment

IEC 61010-2-010: 2005 Safety requirements for electrical equipment for measurement, control and

laboratory use - Part 2-010: Particular requirements for laboratory

equipment for the heating of materials

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

requirements - Part 1: General requirements

EN 61326-2-6:2006 Electrical equipment for measurement, control and laboratory use - EMC

requirements - Part 2-6: Particular requirements - In vitro diagnostic (IVD)

medical equipment

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

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