

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

Date: 07.02.2025

LETTER OF AUTHORIZATION

To whom it may concern,

We, Shenzhen Mindray Bio-Medical Electronics Co., Ltd., ("Mindray") manufacturer of biochemical, immunological and coagulation analyzers, reagents and consumables ("Product(s)"), hereby certify that "Echipamed-Plus" SRL, with business office at str. Valea Trandafirilor, 24B, of. 2-7, MD-2001, Chisinau, Republic of Moldova ("You") is our official distributor and local representative for registration, sales and service of the Product(s) in Republic of Moldova ("Territory").

As the manufacturer, Mindray guarantees the Product(s) 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, 2025**. 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 Product(s), nor allow any party to seek compensation for goodwill developed during the term of Letter of Authorization or any further extension.

Best regards,


Gao Xiufu


Regional Sales & Marketing Manager, IVD Sales & Marketing Department, Central Asia
Shenzhen Mindray Bio-Medical Electronics Co., Ltd.

**SHENZHEN MINDRAY
BIO-MEDICAL ELECTRONICS CO., LTD.**
Mindray Building, Keji 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





CERTIFICATE

No. QS5 044751 0140 Rev. 05

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.:

SH2305501

Effective Date:

2023-07-01

Expiry Date:

2026-06-30

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Date of Issue: 2023-05-25

(Renee Walker)
Director, US Certification Body, MHS

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CERTIFICATE

No. QS5 044751 0140 Rev. 05

Overall Scope Statement:

Design and Development, Production and Distribution of
 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,
 Endoscope Light Source and Accessories, 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, Coagulation Analyzer and
 Accessories, Coagulation Reagents, Calibrators and
 Controls for Coagulation Analyzer, Automated Digital Cell
 Morphology Analyzer, Ion-Selective Electrodes,
 Disposable Anesthesia Mask, Reusable Anesthesia Mask,
 Respiratory Mask, Disposable Breathing Circuit, Reusable
 Breathing Circuit, Heat and Moisture Exchanger, Filter,
 Breathing Bag, Tympanic Thermometer, Wireless Module,
 Wireless Device

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Date of Issue: 2023-05-25

(Renee Walker)
 Director, US Certification Body, MHS

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TUV®





America

CERTIFICATE

No. QS5 044751 0140 Rev. 05

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 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, Endoscope Light Source and Accessories, 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, Coagulation Analyzer and Accessories, Coagulation Reagents, Calibrators and Controls for Coagulation Analyzer, Automated Digital Cell Morphology Analyzer, Ion-Selective Electrodes, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag, Tympanic Thermometer, Wireless Module, Wireless Device

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Date of Issue: 2023-05-25

(Renee Walker)
Director, US Certification Body, MHS

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No. QS5 044751 0140 Rev. 05

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

Design and Development, Production and Distribution of 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, Endoscope Light Source and Accessories, 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, Coagulation Analyzer and Accessories, Coagulation Reagents, Calibrators and Controls for Coagulation Analyzer, Automated Digital Cell Morphology Analyzer, Ion-Selective Electrodes, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag, Tympanic Thermometer, Wireless Module, Wireless Device

(Renee Walker)
Director, US Certification Body, MHS





DAkkS

Deutsche
Akkreditierungsstelle
D-ZM-11321-01-00

Product Service

Certificate

No. Q5 044751 0164 Rev. 05

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). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 044751 0164 Rev. 05

Report No.: SH2305501

Valid from: 2023-09-01
Valid until: 2026-08-31

Date, 2023-06-19

Christoph Dicks
Head of Certification/Notified Body





DAkkS

Deutsche
Akkreditierungsstelle
D-ZM-11321-01-00

Product Service

Certificate

No. Q5 044751 0164 Rev. 05

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

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)

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

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)



Certificate

No. Q5 044751 0164 Rev. 05

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, Endoscope Light Source and accessories, 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, Coagulation Analyzer and Accessories, Coagulation Reagents, Calibrators and Controls for Coagulation Analyzer, Automated Digital Cell Morphology Analyzer, Ion-Selective Electrodes, Disposable Anesthesia Mask, Reusable Anesthesia Mask, Respiratory Mask, Disposable Breathing Circuit, Reusable Breathing Circuit, Heat and Moisture Exchanger, Filter, Breathing Bag, Tympanic Thermometer.



Declaration of Conformity



Manufacturer: Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
Mindray Building, Keji 12th Road South, High-Tech Industrial
Park, Nanshan, 518057, Shenzhen, P. R. China

Manufacturer SRN: CN-MF-000014156

Authorized Representative: Shanghai International Holding Corp. GmbH (Europe)
Eiffestraße 80 20537 Hamburg, Germany

Product: Chemistry Analyzer

Model: BS-430、BS-450、BS-460、BS-410、BS-410E、BS-410S、
BS-470、BS-470E

Basic UDI-DI: 69449040SHYQ-BA43*****N9

Intended Purpose: The system is an chemistry analyzer for in vitro diagnostic use in
clinical laboratories and designed for in vitro quantitative
determination of clinical chemistries in serum, plasma, urine or
cerebrospinal fluid samples.

Classification: Class A (According to Rule 5 of IVDR annex VIII)

Conformity Assessment Route: Annex II and III of IVDR

GMDN code: 56676

We declare that the above mentioned products meet the provisions of the
**REGULATION (EU) 2017/746 OF THE EUROPEAN PARLIAMENT and OF THE
COUNCIL**. All supporting documentations are retained under the premises of the
manufacturer. This declaration of conformity is issued under the sole responsibility of
the manufacturer.

References to CS: /

Notified Body: /

Notified Body No. : /

Identification of the Certificate: /

Start of CE-Marking: 2022.4.25

I hereby am appointed as the authorized person to deal with all the registration and quality
management affairs in my capacity as Manager of Technical Regulation Department of Shenzhen
Mindray Bio-Medical Electronics Co., Ltd, Effective immediately.

Place, Date of Issue:

Shenzhen, 2022.4.25

Signature:



Name of Authorized Signatory:

Mr. Wang Xinbing

Position Held in Company:

Deputy Director, Technical Regulation Department

Applied Standards List

Product: Chemistry Analyzer

Model: BS-430、BS-450、BS-460、BS-410、BS-410E、BS-410S、BS-470、BS-470E

Standards Applied:

EN ISO 18113-1:2011	In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 1: Terms, definitions and general requirements (ISO 18113-1:2009)
EN ISO 18113-3:2011	In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 3: In vitro diagnostic instruments for professional use (ISO 18113-3:2009)
EN ISO 15223-1:2016	Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements
EN ISO 14971:2019	Medical devices - Application of risk management to medical devices (ISO 14971:2019)
EN 13612:2002/AC:2002	Performance evaluation of in vitro diagnostic medical devices
EN 62304:2006/A1:2015	Medical device software - Software life-cycle processes
EN 61010-1:2010/A1:2019	Safety requirements for electrical equipment for measurement, control, and laboratory use Part 1: General requirement
EN IEC 61010-2-081:2020	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
IEC 61010-2-101:2018	Safety requirements for electrical equipment for measurement, control and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

EN 61010-2-010:2020	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 62366-1:2015	Medical devices - Part 1: Application of usability engineering to medical devices
EN 61326-1:2013	Electrical equipment for measurement, control and laboratory use - EMC requirements - Part 1: General requirements
EN 61326-2-6:2013	Electrical equipment for measurement, control and laboratory use - EMC requirements - Part 2-6: Particular requirements - In vitro diagnostic (IVD) medical equipment
BS ISO 20916:2019	In vitro diagnostic medical devices — Clinical performance studies using specimens from human subjects — Good study practice

Declaration of Conformity –V1.0

Declaration of Conformity

Manufacturer: Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
Mindray Building, Keji 12th Road South, High-Tech Industrial
Park, Nanshan, 518057, Shenzhen, P. R. China

EC-Representative: Shanghai International Holding Corp. GmbH (Europe)
Eiffestraße 80
20537 Hamburg, Germany

Product: Chemistry Analyzer

Model: BS-430、BS-450、BS-460、BS-410、BS-410E、BS-410S、
BS-470、BS-470E

We herewith declare under our sole responsibility that the above mentioned products meet the provisions of the Council Directive 2011/65/EU. All supporting documentations are retained under the premises of the manufacturer.

Standards Applied:

EN IEC 63000: 2018 Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances

Start of CE-Marking: 2022.4.25

Place, Date of Issue: Shenzhen. 2022.4.25

Signature:



Name of Authorized Signatory: Mr.WangXinBing

Position Held in Company: Deputy Director, Technical Regulation Department

Declaration of Conformity



Manufacturer: Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
Mindray Building, Keji 12th Road South, High-Tech
Industrial Park, Nanshan, 518057, Shenzhen, P. R. China

Manufacturer SRN: CN-MF-000014156

Authorized Representative: Shanghai International Holding Corp. GmbH (Europe)
Eiffestraße 80 20537 Hamburg, Germany

Product: See Attachment I

Catalogue Number: See Attachment I

Classification: See Attachment I

Conformity Assessment Route: Annex IX excluding CHAPTER II

We declare that the above mentioned products meet the provisions of the REGULATION (EU) 2017/746 OF THE EUROPEAN PARLIAMENT and OF THE COUNCIL. All supporting documentations are retained under the premises of the manufacturer. This declaration of conformity is issued under the sole responsibility of the manufacturer.

References to CS: /

Notified Body: TÜV SÜD Product Service GmbH Ridlerstraße 65
80339 München, Germany.

Notified Body No. : 0123

Identification of the Certificate: NO. V12 044751 0190

Start of CE-Marking: 2022.8.25

I hereby am appointed as the authorized person to deal with all the registration and quality management affairs in my capacity as Manager of Technical Regulation Department of Shenzhen Mindray Bio-Medical Electronics Co., Ltd, Effective immediately.

Place, Date of Issue: Shenzhen, 2024.12.11

Signature:

Name of Authorized Signatory:

Position Held in Company:



Bobby Liu

Manager, Technical Regulation Department



Attachment I

NO	Product Name	Catalogue Number	Classification
1	Lipoprotein (a) Kit (Latex Immunoturbidimetric Method)	105-019437-00	Class B (According to Rule 6 of IVDR Annex VIII)
2		105-019439-00	
3		105-019441-00	
4	Lipoprotein (a) Control	105-019443-00	Class B (According to Rule 6 of IVDR Annex VIII)
5	Total Cholesterol Kit (CHOD-POD Method)	105-000820-00	Class B (According to Rule 6 of IVDR Annex VIII)
6		105-000859-00	
7		105-001595-00	
8	Triglycerides Kit (GPO-POD Method)	105-000821-00	Class B (According to Rule 6 of IVDR Annex VIII)
9		105-000860-00	
10		105-001596-00	
11	Alkaline Phosphatase Kit (IFCC Modified Method)	105-000816-00	Class B (According to Rule 6 of IVDR Annex VIII)
12		105-000855-00	
13		105-004593-00	
14	Albumin Kit (Bromocresol Green Method)	105-000822-00	Class B (According to Rule 6 of IVDR Annex VIII)
15		105-000861-00	
16		105-001597-00	
17	α -Amylase Kit (IFCC Method)	105-000847-00	Class C (According to Rule 3 of IVDR Annex VIII)
18		105-000886-00	
19	Bilirubin Direct Kit (VOX Method)	105-000827-00	Class C (According to Rule 3 of IVDR Annex VIII)
20		105-000866-00	
21		105-004599-00	
22	Bilirubin Total Kit (VOX Method)	105-000826-00	Class C (According to Rule 3 of IVDR Annex VIII)
23		105-000865-00	
24		105-004598-00	
25	HbA1c Calibrator	105-003680-00	Class C (According to Rule 3 of IVDR Annex VIII)
26	HbA1c Control P	105-002138-00	Class C (According to Rule 3 of IVDR Annex VIII)
27	HbA1c Control N	105-002140-00	Class C (According to Rule 3 of IVDR Annex VIII)
28	Uric Acid Kit (Uricase-Peroxidase Method)	105-000887-00	Class B (According to Rule 6 of IVDR Annex VIII)

29	Urea Kit (Urease-GLDH, UV Method)	105-000824-00	Class B (According to Rule 6 of IVDR Annex VIII)
30		105-000863-00	
31		105-004597-00	
32	Phosphorus Kit (Phosphomolybdate Method)	105-000833-00	Class B (According to Rule 6 of IVDR Annex VIII)
33		105-000872-00	
34		105-001608-00	
35	Calcium Kit (Arsenazo III Method)	105-000825-00	Class B (According to Rule 6 of IVDR Annex VIII)
36		105-000864-00	
37		105-001600-00	
38	C-Reactive Protein Kit (Turbidimetry Method)	105-000841-00	Class C (According to Rule 3 of IVDR Annex VIII)
39		105-000880-00	
40		105-004605-00	
41	Rheumatoid Factor Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-002179-00	Class B (According to Rule 6 of IVDR Annex VIII)
42		105-002161-00	
43	RF Calibrator	105-003683-00	Class B (According to Rule 6 of IVDR Annex VIII)
44	Antistreptolysin "O" Kit (Latex Immunoturbidimetric Method)	105-009291-00	Class B (According to Rule 6 of IVDR Annex VIII)
45		105-004630-00	
46		105-004631-00	
47		105-007673-00	
48		105-007674-00	
49		105-007675-00	
50	Antistreptolysin "O" Calibrator	105-004644-00	Class B (According to Rule 6 of IVDR Annex VIII)
51	Alanine Aminotransferase Kit (IFCC Method)	105-000814-00	Class B (According to Rule 6 of IVDR Annex VIII)
52		105-000853-00	
53		105-004591-00	
54	Aspartate Aminotransferase Kit (IFCC Method)	105-000815-00	Class B (According to Rule 6 of IVDR Annex VIII)
55		105-000854-00	
56		105-004592-00	
57	Creatine Kinase Kit (IFCC Method)	105-004615-00	Class B (According to Rule 6 of IVDR Annex VIII)
58		105-000869-00	
59		105-004600-00	
60	Creatine Kinase-MB Kit (IFCC Method)	105-004616-00	Class C (According to Rule 3 of IVDR Annex VIII)
61		105-000870-00	
62		105-004601-00	
63	CK-MB Calibrator	105-001132-00	Class C (According to Rule 3 of IVDR Annex VIII)

64	Glucose Kit (HK Method)	105-000832-00	Class C (According to Rule 3 of IVDR Annex VIII)
65		105-000871-00	
66		105-004609-00	
67	Rheumatoid Factor Kit (Immunoturbidimetric Method)	105-004632-00	Class B (According to Rule 6 of IVDR Annex VIII)
68		105-004633-00	
69		105-004634-00	
70	Rheumatoid Factor Calibrator	105-004645-00	Class B (According to Rule 6 of IVDR Annex VIII)
71	Immunoglobulin A Kit (Turbidimetry Method)	105-004618-00	Class B (According to Rule 6 of IVDR Annex VIII)
72		105-000881-00	
73		105-001617-00	
74	Immunoglobulin M Kit (Turbidimetry Method)	105-000843-00	Class B (According to Rule 6 of IVDR Annex VIII)
75		105-000882-00	
76		105-004606-00	
77	Immunoglobulin G Kit (Turbidimetry Method)	105-004619-00	Class B (According to Rule 6 of IVDR Annex VIII)
78		105-000883-00	
79		105-001619-00	
80	HDL-Cholesterol Kit (Direct Method)	105-000835-00	Class B (According to Rule 6 of IVDR Annex VIII)
81		105-000874-00	
82		105-004610-00	
83	LDL-Cholesterol Kit(Direct Method)	105-000836-00	Class B (According to Rule 6 of IVDR Annex VIII)
84		105-000875-00	
85		105-004611-00	
86	Gamma-Glutamyltransferase Kit (Szasz Method/IFCC stand)	105-000817-00	Class B (According to Rule 6 of IVDR Annex VIII)
87		105-000856-00	
88		105-004594-00	
89	Creatinine Kit (Sarcosine Oxidase Method)	105-004614-00	Class B (According to Rule 6 of IVDR Annex VIII)
90		105-000868-00	
91		105-004612-00	
92	Total Protein Kit (Biuret Method)	105-000823-00	Class B (According to Rule 6 of IVDR Annex VIII)
93	Total Protein in Urine/CSF(TPUC)Kit (Pyrogallol Red-Molybdate Method)	105-009168-00	Class B (According to Rule 6 of IVDR Annex VIII)
94		105-009169-00	
95		105-009170-00	
96	TPUC Control	105-009193-00	Class B (According to Rule 6 of IVDR Annex VIII)
97	High Sensitivity C-reaction Protein Kit (Particle-enhanced)	105-001942-00	Class C (According to Rule 3 of IVDR Annex VIII)
98		105-001943-00	
99		105-001944-00	

	Immunoturbidimetric Assay Method)		
100	HS-CRP Calibrator	105-003685-00	Class C (According to Rule 3 of IVDR Annex VIII)
101	Urea Kit (Urease-GLDH, UV Method)	105-000818-00	Class B (According to Rule 6 of IVDR Annex VIII)
102		105-000857-00	
103		105-004595-00	
104	Transferrin Kit (Immunoturbidimetric Assay Method)	105-004507-00	Class B (According to Rule 6 of IVDR Annex VIII)
105		105-006178-00	
106		105-006177-00	
107		105-002246-00	
108		105-004508-00	
109		105-002247-00	
110	TRF Calibrator	105-002317-00	Class B (According to Rule 6 of IVDR Annex VIII)
111	Iron (Fe) Kit (Colorimetric Assay)	105-002198-00	Class B (According to Rule 6 of IVDR Annex VIII)
112		105-002199-00	
113	Carbon Dioxide (CO2) Kit (Enzymatic Method)	105-002190-00	Class B (According to Rule 6 of IVDR Annex VIII)
114		105-002191-00	
115	Complement C3 Kit (Turbidimetry Method)	105-004617-00	Class B (According to Rule 6 of IVDR Annex VIII)
116		105-000878-00	
117		105-001614-00	
118	Complement C4 Kit (Turbidimetry Method)	105-000840-00	Class B (According to Rule 6 of IVDR Annex VIII)
119		105-000879-00	
120		105-004604-00	
121	Apolipoprotein A1 Kit (Turbidimetry Method)	105-000837-00	Class B (According to Rule 6 of IVDR Annex VIII)
122		105-000876-00	
123		105-004602-00	
124	Apolipoprotein B Kit (Turbidimetry Method)	105-000838-00	Class B (According to Rule 6 of IVDR Annex VIII)
125		105-000877-00	
126		105-004603-00	
127	Ferritin Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-006175-00	Class C (According to Rule 3 of IVDR Annex VIII)
128		105-006176-00	
129		105-002244-00	
130		105-002245-00	
131		105-004505-00	
132		105-004506-00	

133	FER Calibrator	105-002311-00	Class C (According to Rule 3 of IVDR Annex VIII)
134	Microalbumin Kit (Immunoturbidimetric Assay Method)	105-006173-00	Class B (According to Rule 6 of IVDR Annex VIII)
135		105-002242-00	
136		105-002243-00	
137		105-006174-00	
138		105-004503-00	
139		105-004504-00	
140	MALB Calibrator	105-002315-00	Class B (According to Rule 6 of IVDR Annex VIII)
141	MALB Control	105-002316-00	Class B (According to Rule 6 of IVDR Annex VIII)
142	α -Hydroxybutyrate Dehydrogenase Kit (DGKC Method)	105-000819-00	Class B (According to Rule 6 of IVDR Annex VIII)
143		105-000858-00	
144		105-004596-00	
145	Total Bile Acids Kit (Enzymatic Cycling Assay)	105-000867-00	Class B (According to Rule 6 of IVDR Annex VIII)
146		105-001603-00	
147		105-004613-00	
148	Lipase Kit (Enzymatic Colorimetric Assay Method)	105-002171-00	Class B (According to Rule 6 of IVDR Annex VIII)
149		105-002172-00	
150		105-002173-00	
151	Fructosamine (FUN) Kit (Colorimetric Assay)	105-002195-00	Class B (According to Rule 6 of IVDR Annex VIII)
152		105-002196-00	
153	FUN Control	105-020477-00	Class B (According to Rule 6 of IVDR Annex VIII)
154	Immunoglobulin E Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-020854-00	Class B (According to Rule 6 of IVDR Annex VIII)
155		105-004501-00	
156		105-004502-00	
157		105-020853-00	
158		105-002240-00	
159		105-002241-00	
160	IgE Calibrator	105-002309-00	Class B (According to Rule 6 of IVDR Annex VIII)
161	D-Dimer Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-012738-00	Class C (According to Rule 3 of IVDR Annex VIII)
162		105-002236-00	
163		105-002237-00	
164		105-012737-00	

165		105-004497-00	
166		105-004498-00	
167	D-Dimer Calibrator	105-002300-00	Class C (According to Rule 3 of IVDR Annex VIII)
168	D-Dimer Control	105-002301-00	Class C (According to Rule 3 of IVDR Annex VIII)
169	Homocysteine (HCY) Kit (Enzymatic Cycling Method)	105-009174-00	Class B (According to Rule 6 of IVDR Annex VIII)
170		105-009175-00	
171		105-009176-00	
172	HCY Control	105-009194-00	Class B (According to Rule 6 of IVDR Annex VIII)
173	Adenosine Deaminase Kit (Enzymatic Colorimetric Method)	105-003177-00	Class B (According to Rule 6 of IVDR Annex VIII)
174		105-003120-00	
175		105-003125-00	
176		105-026284-00	
177		105-026285-00	
178		105-026286-00	
179	ADA Calibrator	105-003687-00	Class B (According to Rule 6 of IVDR Annex VIII)
180	ADA Control	105-020473-00	Class B (According to Rule 6 of IVDR Annex VIII)
181	Unsaturated Iron Binding Capacity Kit (Colorimetric Method)	105-009265-00	Class B (According to Rule 6 of IVDR Annex VIII)
182		105-004515-00	
183		105-004516-00	
184	UIBC Calibrator	105-002306-00	Class B (According to Rule 6 of IVDR Annex VIII)
185	Retinol Binding Protein Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-009269-00	Class B (According to Rule 6 of IVDR Annex VIII)
186		105-002250-00	
187		105-002251-00	
188		105-006182-00	
189		105-004511-00	
190		105-004512-00	
191	RBP Calibrator	105-002304-00	Class B (According to Rule 6 of IVDR Annex VIII)

192	RBP Control	105-002305-00	Class B (According to Rule 6 of IVDR Annex VIII)
193	Angiotensin Converting Enzyme Kit (Enzymatic Colorimetric Assay Method)	105-006179-00	Class B (According to Rule 6 of IVDR Annex VIII)
194		105-002248-00	
195		105-002249-00	
196		105-006180-00	
197		105-004509-00	
198		105-004510-00	
199	ACE Calibrator	105-002313-00	Class B (According to Rule 6 of IVDR Annex VIII)
200	ACE Control	105-002314-00	Class B (According to Rule 6 of IVDR Annex VIII)
201	5'-Nucleotidase Kit (Enzymatic Colorimetric Method)	105-003119-00	Class B (According to Rule 6 of IVDR Annex VIII)
202		105-003124-00	
203		105-026281-00	
204		105-026282-00	
205		105-026283-00	
206	5'-NT Calibrator	105-003688-00	Class B (According to Rule 6 of IVDR Annex VIII)
207	5'-NT Control	105-020475-00	Class B (According to Rule 6 of IVDR Annex VIII)
208	Glucose-6-Phosphate Dehydrogenase Kit (UV Enzymatic Method)	105-009264-00	Class C (According to Rule 3 of IVDR Annex VIII)
209		105-002254-00	
210		105-002255-00	
211	G6PD Control	105-002308-00	Class C (According to Rule 3 of IVDR Annex VIII)
212	β -Hydroxybutyrate Kit (Enzymatic Colorimetric Method)	105-006184-00	Class B (According to Rule 6 of IVDR Annex VIII)
213		105-004513-00	
214		105-004514-00	
215	β -HB Calibrator	105-002319-00	Class B (According to Rule 6 of IVDR Annex VIII)
216	β -HB Control	105-002320-00	Class B (According to Rule 6 of IVDR Annex VIII)
217		105-003123-00	

218	α -L-Fucosidase Kit (CNPF Method)	105-003128-00	Class C (According to Rule 3 of IVDR Annex VIII)
219	AFU Control	105-020474-00	Class C (According to Rule 3 of IVDR Annex VIII)
220	Cholinesterase (CHE) Kit (DGKC Method)	105-002162-00	Class B (According to Rule 6 of IVDR Annex VIII)
221		105-002163-00	
222	Cystatin C Kit (Latex Immunoturbidimetric Method)	105-004638-00	Class B (According to Rule 6 of IVDR Annex VIII)
223		105-004639-00	
224		105-004640-00	
225	Cystatin C Calibrator	105-004647-00	Class B (According to Rule 6 of IVDR Annex VIII)
226	Cystatin C Control	105-004651-00	Class B (According to Rule 6 of IVDR Annex VIII)
227	Myoglobin Kit (Particle-enhanced Immunoturbidimetric Assay Method)	105-012736-00	Class C (According to Rule 3 of IVDR Annex VIII)
228		105-002238-00	
229		105-002239-00	
230		105-012735-00	
231		105-004499-00	
232		105-004500-00	
233	MYO Calibrator	105-002302-00	Class C (According to Rule 3 of IVDR Annex VIII)
234	Prealbumin Kit (Turbidimetry Method)	105-000845-00	Class B (According to Rule 6 of IVDR Annex VIII)
235		105-000884-00	
236		105-004607-00	
237	Prealbumin Calibrator	105-001130-00	Class B (According to Rule 6 of IVDR Annex VIII)
238	Glucose Kit (GOD-POD Method)	105-000888-00	Class C (According to Rule 3 of IVDR Annex VIII)
239	β 2-Microglobulin Kit (Latex Immunoturbidimetric Method)	105-004641-00	Class B (According to Rule 6 of IVDR Annex VIII)
240		105-004642-00	
241		105-004643-00	
242	β 2-Microglobulin Calibrator(for Serum)	105-004648-00	Class B (According to Rule 6 of IVDR Annex VIII)

243	β 2-Microglobulin Calibrator(for Urine)	105-004649-00	Class B (According to Rule 6 of IVDR Annex VIII)
244	β 2-Microglobulin Control	105-004652-00	Class B (According to Rule 6 of IVDR Annex VIII)
245	Multi Sera Calibrator	105-001144-00	Class C (According to Rule 3 of IVDR Annex VIII)
246	Specific Proteins Calibrator	105-001129-00	Class C (According to Rule 3 of IVDR Annex VIII)
247	Lipids Calibrator	105-001128-00	Class B (According to Rule 6 of IVDR Annex VIII)
248	Multimmun control	105-002303-00	Class C (According to Rule 3 of IVDR Annex VIII)
249	ClinChem Multi Control (level 1)	105-009119-00	Class C (According to Rule 3 of IVDR Annex VIII)
250	ClinChem Multi Control (level 2)	105-009120-00	Class C (According to Rule 3 of IVDR Annex VIII)
251	ASO/CRP/RF Triple Control	105-004650-00	Class C (According to Rule 3 of IVDR Annex VIII)
252	CO2 and TBA Multi Control	105-020476-00	Class B (According to Rule 6 of IVDR Annex VIII)
253	Hemoglobin A1c Kit (Enzymatic Assay Method)	105-009338-00	Class C (According to Rule 3 of IVDR Annex VIII)
254		105-002167-00	
255		105-005738-00	