

Certificate Identification:

SC-09H61

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
09Н61-01	61165	CELL-DYN Emerald 22 LYSE	Self-declared

Authorized European	ABBOTT
Representative	Max-Planck-Ring-2
(Name and Address)	65205 Wiesbaden, Germany
Storage site of technical	Abbott Laboratories
documentation	4551 Great America Parkway
(Name and Address)	Santa Clara, CA 95054
	BIT Group France
	Parc Euromedecine II,
	Rue de la Valsiere
	34 099 – Montpellier, Cedex 5 France
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Signature:

Mirna DiPano

Full Name:

Kevin Richardson

Full Name: Position:

Director of Regulatory Affairs

Position:

Manager, Supplier Quality

Date of Approval:

10-50/4-2017

Date of Approval:

Date Issued:

JUL 1 0 2017

Place Issued:

Abbott Santa Clara

Supersedes:

IRIS V1, April 15, 2016

Effective (Date or Lot Number):

JUL 10 2017



Certificate Identification:

SC-09H62

Legal Manufacturer's Name:

**Abbott Laboratories Diagnostics Division** 

Legal Manufacturer's Address:

Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
09Н62-01	58237	CELL-DYN Emerald 22 DILUENT	Self-declared

Authorized European	ABBOTT	
Representative	Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation	4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054	
	Avantor Performance Materials B.V. Teugseweg 20 Deventer, Overijssel Netherlands 7418 AM  Avantor Performance Materials Poland S.A. ul. Sowinskiego 11 44-101 Gliwice, Poland	
Harmonized Standards	Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:	levi

Signature:

Mirna DiPano

Full Name:

Kevin Richardson

Full Name:

Position:

Manager, Supplier Quality

Position:

Director of Regulatory Affairs

Date of Approval:

Date of Approval:

Date Issued:

JUL **10** 2017

Place Issued:

Abbott Santa Clara

Supersedes:

IRI S V1, April 15, 2016

Effective (Date or Lot Number):

JUL 10 2017







## Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 9001:2015

This is to certify that:

Abbott Laboratories Diagnostics Division

100 Abbott Park Road

Abbott Park Illinois 60064 USA

Holds Certificate Number: FM 743464

and operates a Quality Management System which complies with the requirements of ISO 9001:2015 for the following scope:

Design, Manufacture, Development, Installation, Service and Support of In Vitro Diagnostic Products including Test Kits, Reagents, Accessories and Instruments.

For and on behalf of BSI:

Andrew Launn, EMEA Systems Certification Director

Original Registration Date: 2018-10-12

Latest Revision Date: 2021-10-12

Effective Date: 2021-10-13 Expiry Date: 2022-04-12

Page: 1 of 2





...making excellence a habit.™

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract. An electronic certificate can be authenticated <u>online</u>. Printed copies can be validated at www.bsigroup.com/ClientDirectory

Certificate No: FM 743464

Illinois 60064 USA

Location Registered Activities Abbott Laboratories Diagnostics Division Design, Manufacture, Development, Installation, Service and Support of In Vitro Diagnostic Products including Test Kits, 100 Abbott Park Road Reagents, Accessories and Instruments. Abbott Park Illinois 60064 **USA** Oversight of the Quality Management System for the Abbott Abbott Laboratories Diagnostics Division **Diagnostics Division Sites** - Conway Park 675 North Field Drive Lake Forest Illinois 60045 **USA** Distribution of In Vitro Diagnostic Products including Test Abbott Laboratories Diagnostics Division Kits, Reagents, Accessories and Instruments. - K Complex - Distribution Center Route 41 & Martin Luther King Drive North Chicago

Original Registration Date: 2018-10-12 Effective Date: 2021-10-13 Latest Revision Date: 2021-10-12 Expiry Date: 2022-04-12

Page: 2 of 2

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract. An electronic certificate can be authenticated <u>online</u>. Printed copies can be validated at www.bsigroup.com/ClientDirectory



Certificate Identification: SC-09H72

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address: Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
09H72-01	55866	CELL-DYN 22 Plus Control, Full Pack	Self-declared
09H72-02	55866	CELL-DYN 22 Plus Control, Half Pack	Self-declared

Authorized European	ABBOTT
Representative Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany
Storage site of technical	Abbott Laboratories
documentation 4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054
	Streck, Inc.
	7002 S. 109 <sup>th</sup> Street
	La Vista, NE 68128 USA
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:	M wan	Signature:	Car.
Full Name:	Alfred Evans	Full Name:	Thao Phan
Position:	Director, Quality Assurance	Position:	Associate Director, Regulatory Affairs
Date of Approval:	May 9, 2019	Date of Approval:	May 9, 2019
Date Issued:	MAY 0 0 2019	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V1 April 15, 2016	Effective (Date or Lot Number):	MAY 0 9 2019



**Certificate Identification:** 

SC-09H60-

Legal Manufacturer's Name:

Abbot Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
09H60-01	58236	CELL-DYN Emerald 22 Easy Cleaner	Self-Declared
09Н60-03	58236	CELL-DYN Emerald 22 Easy Cleaner	Self-Declared

Authorized European Representative	Abbott GmbH
(name and address)	Max-Planck-Ring-2
	65205 Wiesbaden, Germany
Storage site of technical	Abbott Laboratories
documentation (name and address)	4551 Great America Parkway
	Santa Clara, CA 95054 USA
	Avantor Performance Materials Poland, S.A
	ul. Sowinskiego 11
	44-101 Gliwice, Poland
Harmonized Standards	Live 1: 4 That is 1D
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices, and Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and Annex VI of the ROHS Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Signature:

Full Name:

Cheryl Nowlan

Full Name:

Thao Phan

Position:

Director, Quality Assurance

Associate Director, Regulatory

Position: **Affairs** 

12 OCT 2020

Date of Approval:

Date of Approval:

Date of Issue:

OCT 12 2020

Place Issued:

Abbott Santa Clara

Supersedes:

September 24, 2020

Effective (Date or Lot Number)

OCT 12 2020



# **CERTIFICATE**OF REGISTRATION

This is to certify that the management system of:

## **Medica Corporation**

(FIN F002402)

Main Site: 5 Oak Park Drive, Bedford, Massachusetts, 01730, United States

Additional Site: 3 Oak Park Drive, Bedford, Massachusetts, 01730, United States

has been registered by Intertek, an MDSAP recognized auditing organization, as conforming to the requirements of:

### ISO 13485:2016

**Brazil**: Federal Law n. 6360/76; RDC ANVISA n. 16/2013; RDC ANVISA n. 23/2012; RDC ANVISA n. 67/2009; RDC ANVISA n. 56/2001

Canada: Medical Devices Regulations – Part 1- SOR 98/282

United States: 21 CFR 820, 21 CFR 803, 21 CFR 806, 21 CFR 807 (Subparts A to D)

Japan: MHLW Ministerial Ordinance 169, Article 4 to Article 68; PMD Act

### The management system is applicable to:

Design, Development, Manufacture, Service, Installation and Distribution of in-vitro diagnostic medical devices, in-vitro diagnostic test kits, in-vitro diagnostic reagents, in-vitro diagnostic analyzers/software used in diagnosis and management of cancer, immune status, disease status, autoimmune status, cardiac markers, protein metabolism, endocrine disorders, blood analytes, urinalysis, blood gases.

**Certificate Number:** 

0089217

**Initial Certification Date:** 

2019-04-19

**Certification Effective Date:** 

2019-04-19

**Certification Expiry Date:** 

2022-04-18





President, Business Assurance

Intertek Testing Services NA, Inc. 900 Chelmsford Street Lowell, MA, USA 01851







Medica Corporation 5 Oak Park Drive Bedford, Massachusetts 01730 Tel 781 275 4892 Fax 781 275 2731 www.medicacorp.com

## 

Product Name: Model/Type:

EasyLyte and accessories per attachment EasyLyte Na/K, Na/K/CI, Na/K/CI, Na/K/CI/Li,

Na/K/Ca/pH, Na/K/Cl/Ca/Li

EasyElectrolytes and accessories per attachment EasyElectrolytes Na/K/Cl, Na/K/Li

#### Manufacturer

Medica Corporation5 Oak Park Drive, Bedford, Massachusetts, 01730, USA

### Representative

EC REP Emergo Europe, Prinsessegracht 20,

2514 AP The Hague, The Netherlands

Tel: +31 70 345 8570 Fax: +31 70 346 7299

### **Means of Conformity**

Medica Corporation declares that the products listed are covered by Annex III of Directive 98/79/EC. These products are self-certified since they are for professional use only and are not listed on Annex II, List A or Annex II, List B of Directive 98/79/EC. In addition, they are in conformity with the Annex I, "Essential Requirements" and provisions of council Directive 98/79/EC for In Vitro Diagnostic Medical Devices, Directive 2011/65/EU Restriction of Hazardous Substance in Electrical and Electronic Equipment, and the corresponding national laws of the Member States.

Place and Date: Bedford, Massachusetts, USA, September 27, 2018

Signature:

**Name:** Photios Makris, Ph.D. **Title:** VP, Regulatory Affairs

Photio dabris

### **EasyLyte Accessories**

Catalog No.	Accessory	EDMA Code
2004	EasyLyte Na/K Analyzer	21 07 11 02
2014	EasyLyte Plus Na/K/Cl Analyzer	21 07 11 02
2015	EasyLyte Lithium Na/K/Li Analyzer	21 07 11 02
2016	EasyLyte Calcium Na/K/Ca/pH Analyzer	21 07 11 02
2021	EasyLyte Na/K/Cl/Li Analyzer	21 07 11 02
2030	EasyLyte EXPAND Analyzer, Na/K/Cl/Ca-Li	21 07 11 02
2070	EasyLyte EasySampler	21 07 11 02
2101	EasyLyte K+ Electrode	11 04 01 06
2102	EasyLyte Na+ Electrode	11 04 01 07
2113	EasyLyte Cl- Electrode	11 04 01 03
2106	EasyLyte Li+ Electrode	11 04 01 04
2150	EasyLyte Ca++ Electrode	11 04 01 02
2151	EasyLyte pH Electrode	11 70 31 02
2152	EasyLyte Disposable Reference Electrode	11 04 04 01
2103	EasyLyte Reference Electrode	11 04 04 01
2258	EasyLyte Membrane Assembly	21 07 11 02
2120	EasyLyte Na/K 800 ml Solutions Pack	11 04 04 02
2121	EasyLyte Na/K/Cl 800mL Solutions Pack	11 04 04 02
2122	EasyLyte Na/K/Li 800mL Solutions Pack	11 04 04 02
2123	EasyLyte Na/K/Ca/pH 800mL Solutions Pack	11 04 04 02
2028	EasyLyte Na/K/Cl/Li 400mL Solution Pack	11 04 04 02
2109	EasyLyte Na/K 400mL Solutions Pack	11 04 04 02
2112	EasyLyte Na/K/Cl 400mL Solutions Pack	11 04 04 02
2115	EasyLyte Na/K/Li 400mL Solutions Pack	11 04 04 02
2114	EasyLyte Na/K/Ca/pH 400mL Solutions Pack	11 04 04 02
2026	EasyLyte Na/K/Cl/Li 800mL Solution Pack	11 04 04 02
2124	EasyLyte Na/K/Cl/Ca-Li 800ml Solutions Pack	11 04 04 02
2814	EasyQC Bi-Level Quality Control Kit	11 50 02 04
2815	EasyQC Tri-Level Quality Control Kit	11 50 02 04
2843	EasyLyte Quality Control Sample Cups (60)	21 07 11 02
2118	Daily Cleaning Solution Kit	11 01 01 27
2598	EasyLyte Daily Cleaner Cup	21 07 11 02
2108	EasyLyte Solutions Valve	21 07 11 02
2107	EasyLyte Sample Probe	21 07 11 02
2257	EasyLyte Sample Detector	21 07 11 02

### **EasyLyte Accessories, continued**

Catalog No.	Accessory	<b>EDMA Code</b>
2104	EasyLyte Tubing Kit	21 07 11 02
2100	EasyLyte Calcium Tubing Kit	21 07 11 02
2492	EasyLyte Internal Filling Solution (125mL)	11 04 04 90
2309	EasyLyte Wash Solution (50mL)	11 04 04 90
2111	EasyLyte Urine Diluent (500mL)	11 04 04 90
2577	EasyLyte Standard Solution, Urine (50mL)	11 04 04 90
2323	EasyLyte Probe Wipers (6)	21 07 11 02
2541	EasyLyte Printer Paper (3 rolls)	21 07 11 02
2595	EasyLyte EasySampler Sample Cups, 500uL (500)	21 07 11 02
2596	EasyLyte Sample Cups 2.0mL (500)	21 07 11 02
10745	Anti-Evaporation Caps (500)	21 07 11 02
2293	EasyLyte Capillary Tubes	21 07 11 02
2590	EasyLyte Capillary Adaptor Kit	21 07 11 02
2292	EasyLyte Capillary Adaptor Cleaning Kit	21 07 11 02
2578	EasyLyte Red Dye Test Solution (50mL)	11 30 01 11
2572	EasyLyte Troubleshooting Kit	21 07 11 02
2571	EasyLyte Troubleshooting Kit (Na/K/Ca/pH and Na/K/Cl/Li)	21 07 11 02
2105	EasyLyte Quarterly Operating Kit	21 07 11 02
2095	EasyLyte Maintenace Kit	21 07 11 02
2076	EasyLyte Sample Tray	21 07 11 02
2074	EasyLyte Sample Cup Retainer Ring	21 07 11 02
7118	Daily Rinse/Cleaning Solution Kit	11 01 01 27
2544	EasyLyte C Series Printer Paper (5 rolls)	21 07 11 02
2934	EasyLyte Barcode Reader Kit	21 07 11 02

### **EasyElectrolytes Accessories**

Catalog No.	Accessory	EDMA Code
4002	EasyElectrolyte Na/K/Cl Analyzer	21 07 11 02
4003	EasyElectrolyte Na/K/Li Analyzer	21 07 11 02
4102	Reagent Module, Na/K/Cl	11 04 04 02
4103	Reagent Module, Na/K/Li	11 04 04 02
7205	EasyElectrolyte/EasyStat Na+ Electrode	11 04 01 07
7206	EasyElectrolyte/EasyStat K+ Electrode	11 04 01 06
4203	EasyElectrolyte CI- Electrode	11 04 01 03
4204	EasyElectrolyte Li+ Electrode	11 04 01 04
6204	EasyElectrolyte/EasyStat/EasyBloodGas Reference Electrode	11 04 04 01
4207	EasyElectrolyte Spacer Electrode	11 04 01 90
4301	EasyElectrolyte Troubleshooting Kit	21 07 11 02
2118	Daily Cleaning Solution Kit	11 01 01 27
4402	EasyStat/EasyBloodGas/EasyElectrolyte Red Test Dye Solution	11 30 01 11
4403	EasyElectrolyte Urine Diluent	11 04 04 90
2814	Bi-Level Quality Control Kit	11 50 02 04
2815	Tri-Level Quality Control Kit	11 50 02 04
4405	EasyElectrolyte Na/K/Cl Demonstration Kit	21 07 11 02
4406	EasyElectrolyte Na/K/Li Demonstration Kit	21 07 11 02
4404	EasyElectrolyte Capillary Tube Kit	21 07 11 02
4306	EasyElectrolyte Sampler	21 07 11 02
6504	EasyBloodGas/EasyElectrolyte Pump Tube	21 07 11 02
6505	EasyStat/EasyBloodGas/EasyElectrolyte Printer Paper	21 07 11 02
4506	EasyElectrolyte Sensor Module	21 07 11 02
4507	EasyElectrolyte Valve Module	21 07 11 02
4508	EasyStat/EasyBloodGas/EasyElectrolyte Compression Plate	21 07 11 02
7302	Probe Wipers	21 07 11 02
4522	EasyElectrolyte Daily Cleaner Sample Cups	21 07 11 02
4539	EasyElectrolyte Sensor Module, Li+	21 07 11 02
6537	EasyElectrolyte/EasyStat/EasyBloodGas Serial Cable, 9-pin	21 07 11 02
6520	EasyElectrolyte/EasyStat/EasyBloodGas Barcode Reader Kit	21 07 11 02





## Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016 & EN ISO 13485:2016

This is to certify that: Helena Laboratories (UK) Ltd

trading as Helena Biosciences Europe

**Queensway South** 

**Team Valley Trading Estate** 

Gateshead Tyne and Wear NE11 OSD United Kingdom

Holds Certificate Number: MD 69326

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 & EN ISO 13485:2016 for the following scope:

The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.

For and on behalf of BSI:

Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2002-10-25 Effective Date: 2021-04-14 Latest Revision Date: 2021-04-13 Expiry Date: 2024-04-13

Page: 1 of 2

...making excellence a habit."





This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract. An electronic certificate can be authenticated <u>online</u>.

Printed copies can be validated at www.bsigroup.com/ClientDirectory

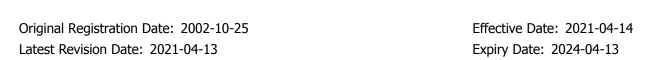
Certificate No: MD 69326

### Location

Helena Laboratories (UK) Ltd trading as Helena Biosciences Europe Sunderland Enterprise Park Colima Avenue Sunderland SR5 3XB United Kingdom Registered Activities

The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.

Helena Laboratories (UK) Ltd trading as Helena Biosciences Europe Queensway South Team Valley Trading Estate Gateshead Tyne and Wear NE11 OSD United Kingdom The design, manufacture, supply, servicing and repair of in-vitro diagnostic devices, molecular biology products, immunochemistry products and medical laboratory equipment and consumables.



Page: 2 of 2

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract. An electronic certificate can be authenticated <u>online</u>.

Printed copies can be validated at www.bsigroup.com/ClientDirectory



HL-7-DC-0814 Rev. 1

In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5560	APTT Si L Minus	55981

I, the undersigned, declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: C.J. Sandercock Title: QA and Regulatory Affairs Officer

Signed: Mandewile Date: 24 Nov 2020

Helena Biosciences Europe, Gateshead, Tyne and Wear, NE11 0SD, United Kingdom Tel +44 (0)191 482 8440 info@helena-biosciences.com www.helena-biosciences.com EC REP

Prince Technologies B.V. Waanderweg 62, 7812 HZ Emmen, The Netherlands



HL-7-0664DC DOI 2015/08 (1)

In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5267L	Thromboplastin L	55983

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson Title: Managing Director

Signed: Michael / Tylen Date: 06 Aug 2015

Tel +44 (0)191 482 8440

Fax +44 (0)191 482 8442

info@helena-biosciences.com

www.helena-biosciences.com

Helena Biosciences Europe Queensway South, Team Valley Trading Estate, Gateshead, Tyne and Wear, NE11 0SD, United Kingdom



HL-7-0512DC DOI 2015/08 (5)

In Application of the Council Directive 98/79/EC on the approximation of the laws of the Member States relating to *In Vitro* Diagnostic Medical Devices & CE marking.

Declaration of conformance to applicable sections of Annex I - Essential Requirements and Annex III (EC Declaration of Conformity) imposed by sections 2 to 5. The below listed products are not classified under Annex II Lists A or B. Access to the appropriate technical files will be made available to the appropriate body in the event this is required.

Product Code	Description	GMDN Classification Code
5556	Clauss Fibrinogen 50	55997

I, the undersigned declare that the devices registered against the above GMDN Classification Code conforms to the said Directives.

Full Name: M.J. Stephenson Title: Managing Director

Signed: Michael / Tylen Date: 12 Aug 2015

Tel+44 (0)191 482 8440Helena Biosciences EuropeFax+44 (0)191 482 8442Queensway South, Team Valley Trading Estate,<br/>Gateshead, Tyne and Wear, NE11 0SD,<br/>United Kingdom



## **CERTIFICATO N° 505SGQ05**

CERTIFICATE N° 505SGQ05

Si certifica che il this is to certify that

### Sistema di Gestione per la Qualità

Quality Management System

messo in atto da

## APTACA S.p.A.

Via Monte Bianco, 4 – IT 20900 MONZA (MB)

nella Sede Operativa di

Regione Monforte, 30 - IT 14053 CANELLI (AT)

è conforme alla norma is in compliance with the standard

UNI EN ISO 9001-2015 (ISO 9001-2015)

per i seguenti Processi concerning the following kinds of Processes

Gestione della fabbricazione ed immissione in commercio di tamponi sterili per il prelievo di campioni biologici in orifizi naturali e in ambito chirurgico. Progettazione e fabbricazione di dispositivi medico diagnostici per laboratori di analisi. Gestione della fabbricazione ed immissione in commercio di dispositivi medici invasivi in relazione agli orifizi del corpo in Classe I Sterile. Fabbricazione di dispositivi medici invasivi in relazione agli orifizi del corpo in Classe I Sterile. Commercializzazione di dispositivi medici e diagnostici in vitro.

#### Commercializzazione di articoli da laboratorio

Management of the manufacturing and placing on the market of sterile tampons for sampling of biological specimens in natural orifice and in surgical field.

Design and manufacturing of diagnostic medical devices for laboratories of analysis. Management of the manufacturing and placing on the market of invasive medical devices with respect to body orifices (class I sterile).

Marketing of medical and diagnostic devices in vitro. Marketing of laboratory articles.

Il presente Certificato è soggetto al rispetto delle condizioni stabilite dai Regolamenti per la certificazione in vigore applicabili.

This Certificate shall satisfy the requirements established in the Rules for the certification in force applicable.

In caso di discordanza tra le lingue utilizzate nella traduzione del contenuto del presente certificato, fare riferimento alla lingua italiana In cases of discrepancy between the languages used in the translation of the content of this certificate, please refer to the Italian language

L'AMMINISTRATORE DELEGATO
MANAGING DIRECTOR

Dr. Ing. Roberto Cusolito

Data di Prima Emissione First Issue Date Data di Prima Emissione ITALCERT

First Issue Date ITALCERT

Data di Rinnovo Renewal Date Data di Scadenza Expiration Date

1998-07-23

2011-10-30

2020-10-30

2023-10-29

Settore IAF 14 - 29



SGQ Nº 023A

Membro degli Accordi di Mutuo Riconoscimento EA, IAF e ILAC

Signatory of EA, IAF and ILAC Mutual Recognition Agreements



# MANAGEMENT SYSTEM CERTIFICATE

Certificate no.: 59878-2009-AQ-MCW-FINAS

Initial certification date:

Valid: 01 September 2021 – 31 August 2024

This is to certify that the management system of

### THERMO FISHER SCIENTIFIC

Kubinskaya 73, liter A, build.1, Saint-Petersburg, Russian Federation, 196240

has been found to conform to the Quality Management System standard:

ISO 9001:2015

This certificate is valid for the following scope:

MANUFACTURING OF LIQUID HANDLING PRODUCTS AND SPECIAL DIAGNOSTIC PLASTICS.

Place and date: Espoo, 18 June 2021







For the issuing office: DNV - Business Assurance Keilaranta 1, 02150 Espoo, Finland

22

Kimmo Haarala Management Representative





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

### CERTIFICATO n. CERTIFICATE No.

4264/4

SI CERTIFICA CHE IL SISTEMA DI GESTIONE PER LA QUALITÀ DI WE HEREBY CERTIFY THAT THE QUALITY MANAGEMENT SYSTEM OPERATED BY

### GRUPPO VACUTEST KIMA

Sede / Head Office

Via dell'Industria, 12 - 35020 Arzergrande (PD) - Italia Unità Operative / Operative Units

MEUS S.r.I. - Via Leonardo da Vinci, 24B - 26 - 28 - Zona Industriale Tognana - 35028 Piove di Sacco (PD) - Italia MEUS S.r.I. - Via dell'Industria, 2 - 16 - 35020 Arzergrande (PD) - Italia

ROLL S.R.L. - Via Leonardo Da Vinci, 24A – Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia KIMA S.R.L. - Via Leonardo da Vinci, 22 – Zona Industriale Tognana – 35028 Piove di Sacco (PD) - Italia VACUTEST KIMA S.r.I. - Via dell'Industria,12 – 35020 Arzergrande (PD) – Italia VACUTEST KIMA S.r.I. via L. Da vInci, 22 Zona Industriale Tognana - 35028 Piove di Sacco (PD) - Italia

È CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

### UNI EN ISO 9001:2015

Sistema di Gestione per la Qualità / Quality Management System PER LE SEGUENTI ATTIVITÀ / FOR THE FOLLOWING ACTIVITIES

EA: 14 - 29

Progettazione e produzione di provette con vuoto predeterminato ad uso prelievo ematico, liquidi biologici e urine. Produzione di provette per microprelievi di sangue. Progettazione e produzione di Holders (camicie) per prelievo sottovuoto. Progettazione e produzione di kit diagnostici per l'analisi del sangue e dei liquidi biologici. Progettazione e produzione di terreni di coltura per microbiologia. Progettazione e produzione di aghi e dispositivi sterili per il prelievo ematico. Commercializzazione di prodotti del Gruppo: kit diagnostici, terreni di coltura per microbiologia, articoli in plastica per laboratorio analisi, provette con vuoto predeterminato e aghi sterili. Progettazione e produzione di stampi per articoli in plastica per laboratorio analisi. Stampaggio di materie termoplastiche ad iniezione per articoli medicali.

Design and production of test tubes with predetermined vacuum for collection of haematological samples, biological liquids and urine samples. Production of test tubes for micro-collection of haematological samples. Design and production of Holders for vacuum sampling. Design and production of diagnostic kits for blood and biological liquids analysis. Design and production of culture media for microbiology. Design and production of sterile needles and devices for collection of haematological samples. Trading of the products of the Group: diagnostic kits, culture media for microbiology, plastic disposable labware, test tubes with predetermined vacuum and sterile needles. Design and production of moulds for plastic labware. Injection moulding of thermoplastic materials for medical devices.

Riferirsi alla documentazione del Sistema di Gestione per la Qualità aziendale per l'applicabilità dei requisiti della norma di riferimento.

Refer to the documentation of the Quality Management System for details of application to reference standard requirements.

Il presente certificato è soggetto al rispetto del documento ICIM "Regolamento per la certificazione dei sistemi di gestione" e al relativo Schema specifico. The use and the validity of this certificate shall satisfy the requirements of the ICIM document "Rules for the certification of company management systems" and specific Scheme.

> Per informazioni puntuali e aggiornate circa eventuali variazioni intervenute nello stato della certificazione di cui al presente certificato, si prega di contattare il n° telefonico +39 02 725341 o indirizzo e-mail info@icim.it.
> For timely and updated information about any changes in the certification status referred to in this certificate,
> please contact the number+39 02 725341 or email address info@icim.it.

Data emissione First issue 18/01/2007 Emissione corrente Current issue 18/01/2019

Data di scadenza Expiring date 17/01/2022

M.g., A.

Piazza Don Enrico Mapelli 75 – 20099 Sesto San Giovanni (MI) www.icim.it



www.cisg.com



### **ELITech Clinical Systems**

Zone industrielle 61500 Sées - France

Tél: +33 (0)2 33 81 21 00 Fax: +33 (0)2 22 28 77 51

www.elitechgroup.com



### **DECLARATION DE CONFORMITE CE**

Nous, ELITech Clinical Systems SAS, zone industrielle 61500 SEES France, déclarons sous notre seule responsabilité que les réactifs référencés dans la liste ci-jointe (2 pages), sont conformes aux exigences essentielles des annexes I et III de la Directive Européenne 98/79/CE relative aux dispositifs médicaux de diagnostic *in vitro* et au code de la santé publique.

Ces dispositifs sont classés dans la catégorie « autre dispositif » puisqu'ils n'appartiennent ni à la liste A et liste B de l'annexe II et ni à la classe des autotests.

Cette déclaration est basée sur le contenu de chaque dossier technique et s'appuie sur la certification de notre système qualité selon la norme NF EN ISO 13485 : 2016 (Certification valable jusqu'au 27 juillet 2023).

### **DECLARATION OF EC CONFORMITY**

We, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, hereby certify, under our own responsibility, that the reagents such as listed attached (2 pages), conform to the essential requirements of appendices I and III of European Directive 98/79/EC, relating to in vitro diagnostic medical devices and to the public health code.

These devices are classified in the "other device" category since they do not belong neither to list A or list B of annex II nor to self-testing class.

This declaration is based on the contents of each technical file and is supported by the certification of our quality system according to the standard NF EN ISO 13485 : 2016 (Certification valid until July 27th, 2023).

### DECLARACIÓN CE DE CONFORMIDAD

Nosotros, ELITech Clinical Systems SAS, Zone Industrielle 61500 SEES France, declaramos bajo nuestra única responsabilidad que los reactivos referenciados en la lista adjunta (2 páginas), son conformes con los requisitos esenciales de los anexos I y III de la Directiva Europea 98/79/CE sobre dispositivos médicos para diagnóstico in vitro y el código de salud pública.

Estos dispositivos se clasifican en la categoría "otro dispositivo", ya que no pertenecen a la lista A ni a la lista B del anexo II, tampoco a la clase de autodiagnóstico.

Esta declaración se basa en el contenido de cada expediente técnico y está respaldado por la certificación de nuestro sistema de calidad según la norma NF EN ISO 13485 : 2016 (Certificación válida hasta el 27 de Julio 2023).

Sées, le 12 Mai 2021

Valérie LAMBERT,

**ELITech Clinical Systems SAS** 

Zone Industrielle

61500 SEES - France

Regulatory Affairs Manager Tél.: Responsable de los Asuntos Reglementarios

Responsable des Affaires Réglementaires

Tél.: +33(0)2 33 81 21 00 - Fax: +33(0)2 33 28 77 51

SIRET 318 365 228 00036

Cécile GOUBAULT,

Directeur Général Délégué

Managing Director Directora General

Société par actions simplifiée au capital de 1.688.392,33 € - SIREN : 318 365 228 - RCS ALENCON

REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCIAS	Code GMDI
M	etabolites divers / Miscellaneous metabolites	
ALBUMIN	ALBU-0600/0700/0250/M830	
ALBUMIN ENVOY	ALBU-0850	53597
BILIRUBIN DIRECT 4+1	BIDI-0600/0250	53233
BILIRUBIN TOTAL 4+1 BILIRUBIN TOTAL & DIRECT 4+1	BITO-0600/0250	53229
CREATININE ENVOY	BITD-0600 CRSL-0850	53229/53233
CREATININE JAFFE	CRCO-0600/0700	53250 53251
CREATININE PAP	CRSL-M490	
CREATININE PAP SL	CRSL-0630/0250	53250
DIRECT BILIRUBIN	BIDI-M430	53233
DIRECT BILIRUBIN ENVOY	BIDV-0850	53233
BLUCOSE ENVOY BLUCOSE HK	GPSL-0850	
BLUCOSE HK SL	GHSL-M490 GHSL-0600/0250	53301
SLUCOSE PAP	GPSL-M690	33301
SLUCOSE PAP SL	GPSL-0507/0500/0707/0700/0250/0455/0497	-1
ACTATE	LACT-0100	53342
IICROPROTEIN PLUS	PRTU-0600/0250	53481
HOSPHORUS	PHOS-0600/0230/M430	59123
HOSPHORUS ENVOY	PHOS-0850	59123
OTAL BILIRUBIN	BITO-M430	53229
OTAL BILIRUBIN ENVOY DTAL PROTEIN	BITV-0850	53229
OTAL PROTEIN OTAL PROTEIN ENVOY	PROB-M830	50005
OTAL PROTEIN PLUS	PROB-0650 PROB-0600/0700/0250	53985
REA	URSL-M830	
REA ENVOY	URSL-0850	53587
REA UV SL	URSL-0427/0420/0500/0507/0250/0455	- 33007
RIC ACID	AUML-M830	
RIC ACID ENVOY	AUVD-0850	E0500
RIC ACID MONO SL	AUML-0497/0427/0420/0500/0507/0707/0250	53583
RIC ACID SL	AUSL-0250	
RINE PROTEIN	PRTU-M230	53481
	Enzymes / Enzymes	
P (DEA) SL	PASL-0400/0420/0230	
P ENVOY	PIVD-0850	52928
P IFCC	ALPI-0230	
T ENVOY	ALSL-0850	
.T/GPT	ALSL-M490	52923
T/GPT 4+1 SL	ALSL-0410/0430/0510/0250/0455	
MYLASE	AMSL-M430	9808
MYLASE ENVOY MYLASE SL	AMSL-0850	52940
ST/GOT	AMSL-0390/0400/0230	
ST ENVOY	ASSL-M490 ASVD-0850	52954
ST/GOT 4+1 SL	ASSL-0410/0430/0510/0250/0455	32934
OLINESTERASE	CHES-0053	52971
ENVOY	CKSL-0850	53003
K-MB ENVOY	CMSL-0850	
-MB SL / CKMB	CMSL-0410/0430/0230	52994
NAC	CKSL-M230	53003
NAC SL	CKSL-0410/0430/0230	55005
MMA-GT MMA-GT PLUS SL	GISL-M230	7240000
T ENVOY	GISL-0400/0420/0250	53027
HENVOY	GISL-0850 LLSL-0850	
H IFCC	LLSL-M230	53072
H-L SL	LLSL-0400/0420/0230	- 33072
ASE	LPSL-0250	
ASE ENVOY	LPSL-0850	53108
ASE SL	LPSL-0230	
Electrolyte	es / Oligo-élements / Electrolytes / Trace-elements	Will be and South the
CIUM ARSENAZO		
CIUM ENVOY	CALA-0600/0250/M430 CALA-0850	45789
ORIDE	CALA-0850 CHLO-0600/0250	
N ENVOY	FEFE-0850	60037
N FERENE	FEFE-0230/0600/M230	54758
SNESIUM ENVOY	MAGX-0850	
GNESIUM XB	MGXB-0250/0600/M430	46795
GNESIUM XYLIDYL	MAGX-0230/0600	
	Lipides / Lipids	
DLESTEROL		
DLESTEROL ENVOY	CHSL 0850	53359
DLESTEROL HDL SL 2G	CHSL-0850 HDLL-0230/0380/0390	
DLESTEROL LDL SL 2G	LDLL-0230/0380/0390	53391
DLESTEROL SL	CHSL-0507/0500/0700/0707/0250/0455/0497	53395 53359
CHOLESTEROL	CHDL-0250/0600/M330	
CHOLESTEROL ENVOY	HDLL-0850	53391
CHOLESTEROL	CLDL-0250/M330	50000
CHOLESTEROL ENVOY	LDLL-0850	53395
GLYCERIDES	TGML-M690	
GLYCERIDES ENVOY TGML-0850		53460
	DOT (1) AND (1) AND (1)	53460
GLYCERIDES ENVOY GLYCERIDES MONO SL NEW	TGML-0427/0425/0515/0700/0517/0707/0497	53460



REACTIFS - REAGENTS - REACTIVOS	RÉFÉRENCES - REFERENCIAS	Code GMDN
Contrôles-Cal	librants-Standards / Controls-Calibrators-Standards	
HOLESTEROL HDL 2G CALIBRATOR	HDLL-0011/0041	44696
HOLESTEROL LDL 2G CALIBRATOR	LDLL-0011/0041	41728
HOLESTEROL Standard 200 mg/dL	CHOL-0055	44698
(-MB CONTROL	CKMB-0900	44693
JICAL 2	CALI-0550	47868
ITROL I	CONT-0060	47869
ITROL II	CONT-0160	41818
_UCOSE Standard 100 mg/dL	GLUP-0055 HLCA-0041	47868
DL LDL CALIBRATOR E CONTROL I	ISCT-0046	, , , , , , , , , , , , , , , , , , , ,
E CONTROL II	ISCT-0047	47869
CROPROTEIN PLUS Standard 100 mg/dL	PRTU-0022	53482
RIGLYCERIDES Standard 200 mg/dL	TRIG-0055	44702
REA Standard 50 mg/dL	URUV-0055	53588
RIC ACID Standard 6 mg/dL	ACUR-0055	44704
Paris In the Paris	rotéines spécifiques / Specific proteins	
ITI-STREPTOLYSIN O	ASLO-0250	59055
RP IP	ICRP-0400/M230	53705
RP IP CALIBRATOR SET	ICRP-0043	41838
RP IP CONTROL I	ICRP-0046	41839
RP IP CONTROL II	ICRP-0047	
RP WR	CRPW-0230	53705
RP WR CALIBRATOR SET	CRPW-0043	41838
RP WR CONTROL	CRPW-0045	41839
RP WR ENVOY	CRPW-0850	53705
ERRITIN	IFRT-0230	53718
ERRITIN CALIBRATOR	IFRT-0042	41927 53737
APTOGLOBIN IP	IHAP-0400 HBAC-0240	59090
bA1c	HBAC-0240 HBAC-0043	53315
batc CALIBRATOR SET	HBAC-0049	44435
bA1c CONTROL L + H	IIGA-0400	53760
A IP	IIGG-0400	53787
G IP	IIGM-0400	53795
M IP ALBUMIN IP	IMAL-0400	53475
ALBUMIN IP CALIBRATOR SET	IMAL-0043	53477
ALBUMIN IP CONTROL I	IMAL-0046	
ALBUMIN IP CONTROL II	IMAL-0047	53478
ROSOMUCOID IP	IORO-0400	53606
REALBUMIN IP	IPAL-0400	53957
ROTEIN IP CALIBRATOR SET	IPRO-0043	53593
F CALIBRATOR	IRFA-0042	42230
HEUMATOID FACTOR	IRFA-0230	55111
HEUMATOLOGY CONTROL I	IRCT-0046	47869
HEUMATOLOGY CONTROL II	IRCT-0047	
RANSFERRIN IP	ITRF-0400	59041
	Vitamines/Vitamins	
ITAMIN D	VITD-0250	54476
ITAMIN D CALIBRATOR SET	VITD-0043	54474
ITAMIN D CONTROL SET	VITD-0049	54475
	Solutions pour électrodes selectives d'ions /	
	SE Solutions for ion-selective electrodes	F0000
SE BASELINE SOLUTION ENVOY	ISBA-0850	59238
SE CALIBRATORS	ISCA-0250	52867
SE CALIBRATOR ENVOY	ISCV-0850	50050
SE CLEANER/CONDITIONER	ISCC-0280	59058
SE DILUENT	ISDI-0250	58237
SE DILUENT ENVOY	ISDV-0850	
E REFERENCE SOLUTION	ISRS-0800	59238
E REFERENCE SOLUTION ENVOY  Solutions de la	ISRS-0850 vage pour les équipements ELITech Clinical Systems /	
	olutions for ELITech Clinical Systems Equipments	
Cicuming of	SLHC-5900	59058
OID DOLLITION for ELiTrob Cileiral Custome Application	John Caper	
	SI NA-5900	59058
YSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers	SLNA-5900 SLSV-5905	59058
CID SOLUTION for ELITech Clinical Systems Analyzers EYSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers EYSTEM SOLUTION EXECUTION CONTROL OF ELITECH Clinical Systems Analyzers	SLSY-5905	59058
YSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers YSTEM SOLUTION YSTEM SOLUTION for ELITech Clinical Systems Analyzers	SLSY-5905 SLSY-5900	
YSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers YSTEM SOLUTION YSTEM SOLUTION for ELITech Clinical Systems Analyzers /ASH SOLUTION A	SLSY-5905	58236
YSTEM CLEANING SOLUTION for ELITech Clinical Systems Analyzers YSTEM SOLUTION YSTEM SOLUTION for ELITech Clinical Systems Analyzers (ASH SOLUTION A (ASH SOLUTION B	SLSY-5905 SLSY-5900 SOLA-M163	58236 59058





### **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 60119814 0001

Report No.: 21265422 001

Manufacturer: Macherey-Nagel GmbH & Co. KG

Neumann-Neander-Str. 6-8

52355 Düren Deutschland

**Products:** Products for self-testing

(see attachment for products and sites included)

Replaces Certificate, Registration No.: HL 60076687 0001

**Expiry Date:** 2022-05-28

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: 2017-05-29

Date: 2017-05-29

5. E 13/1/2

rüvRheinlar

Notified Body

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.



Doc. 1/1, Rev. 0

## TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment to Certificate

Registration No.: HL 60119814 0001 Report No.: 21265422 001

Manufacturer: Macherey-Nagel GmbH & Co. KG

Neumann-Neander-Str. 6-8

52355 Düren Deutschland

Products for self-testing:

- Single and multi-parameter disposable test strips for urine analysis
- indicator test strips and papers for measurement of pH in urine

Additional site for warehousing and logistics:

Bahnstr. 120 52355 Düren, Germany

Notified Body

Dipl.-Ing, Sven Hoffmani

Date: 2017-05-29



ФЕДЕРАЛЬНОЕ АГЕНТСТВО ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ ГОСТ Р «EAC AUDIT» РЕГИСТРАЦИОННЫЙ HOMEP POCC RU.32028.04EAC1 ОРГАН ПО СЕРТИФИКАЦИИ ООО «ГОРТЕСТ» РЕГИСТРАЦИОННЫЙ HOMEP POCC RU.32028 ИНН 7717616798 ОГРН 1087746489060 Юридический адрес: 109028, Россия, г. Москва, Серебряническая набережная, д. 27, этаж 4, пом. 1, ком. 17

Телефон: 8 (800) 1000-730, e-mail: info@eacaudit.ru



№ 005032

## СЕРТИФИКАТ СООТВЕТСТВИЯ

Регистрационный номер № 04EAC1.CM.03842

Общество с ограниченной ответственностью «Агат-Мед»

(наименование лица)

105173, Россия, г. Москва, ул. Главная, д. 6, кв. 12

(поридический адрес лица)

143906, Россия, Московская область, г. Балашиха, квартал Щитниково, д. 88А

(фактический адрес лица)

ИНН: 7719187311

ОГРН: 1037739078970

### НАСТОЯЩИЙ СЕРТИФИКАТ УДОСТОВЕРЯЕТ СООТВЕТСТВИЕ

системы менеджмента качества изделий медицинских Общества с ограниченной ответственностью «Агат-Мед» требованиям ГОСТ ISO 13485-2017 (ISO 13485:2016) «Изделия медицинские. Системы менеджмента качества. Системные требования для целей регулирования» применительно к разработке, производству и продаже медицинских изделий для in vitro диагностики: реагентов и наборов реагентов для клинической биохимии, а также калибраторов и контрольных материалов

Лага регистрации: 08-09-2021

Срок действия до: 07-09-2024

Руководитель органа по сертификации:

(подпись)

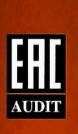
В. И. Погодин

Председатель экспертной комиссии

М.П.

Е. Д. Курбатова

НАСТОЯЩИЙ СЕРТИФИКАТ ОБЯЗЫВАЕТ ОРГАНИЗАЦИЮ ПОДДЕРЖИВАТЬ СОСТОЯНИЕ ВЫПОЛНЯЕМЫХ РАБОТ В СООТВЕТСТВИИ С ВЫШЕУКАЗАННЫМИ СТАНДАРТАМИ, ЧТО БУДЕТ НАХОДИТЬСЯ ПОД КОНТРОЛЕМ ОРГАНА ПО СЕРТИФИКАЦИИ СИСТЕМЫ ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ "EAC AUDIT" И ПОДТВЕРЖДАТЬСЯ ПРИ ПРОХОЖДЕНИИ ЕЖЕГОДНОГО ИНСПЕКЦИОННОГО КОНТРОЛЯ



ФЕДЕРАЛЬНОЕ АГЕНТСТВО
ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ
СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ ГОСТ Р
«EAC AUDIT»
РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028.04EAC1
ОРГАН ПО СЕРТИФИКАЦИИ ООО «ГОРТЕСТ»
РЕГИСТРАЦИОННЫЙ НОМЕР РОСС RU.32028
ИНН 7717616798 ОГРН 1087746489060



Юридический адрес: 109028, Россия, г. Москва, Серебряническая набережная, д. 27,

этаж 4, пом. 1, ком. 17

Телефон: 8 (800) 1000-730, e-mail: info@eacaudit.ru

### **РАЗРЕШЕНИЕ**

### на применение знака соответствия системы добровольной сертификации ГОСТ Р «EAC AUDIT»

Регистрационный номер № 04ЕАС1.СМ.03842

ВЫДАНО НА ОСНОВАНИИ РЕШЕНИЯ О ВЫДАЧЕ СЕРТИФИКАТА СООТВЕТСТВИЯ СИСТЕМЫ МЕНЕДЖМЕНТА КАЧЕСТВА ИЗДЕЛИЙ МЕДИЦИНСКИХ

Общество с ограниченной ответственностью «Агат-Мед»

(наименование лица)

105173, Россия, г. Москва, ул. Главная, д. 6, кв. 12

(юридический адрес лица)

143906, Россия, Московская область, г. Балашиха, квартал Щитниково, д. 88А

(фактический адрес лица)

ИНН: 7719187311

ОГРН: 1037739078970

### **РАЗРЕШАЕТ**

Применять знак соответствия системы добровольной сертификации «ЕАС AUDIT» на период действия сертификата соответствия № 04EAC1.CM.03842 в любой форме, исключающей возможность толкования его как знака соответствия качества продукции. Допускается использовать знак соответствия в рекламных буклетах, проспектах, брошюрах, плакатах, бланках организационно-распорядительной документации организации — держателя сертификата.

Руководитель органа по сертификации:

(подпись)

В. И. Погодин

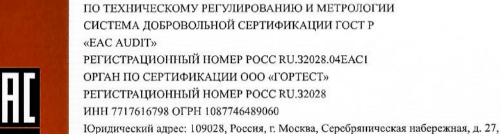
Председатель В экспертной компесии

М.П.

Kypwamokg

Е. Д. Курбатова





ФЕДЕРАЛЬНОЕ АГЕНТСТВО



этаж 4, пом. 1, ком. 17

Телефон: 8 (800) 1000-730, e-mail: info@eacaudit.ru

## СЕРТИФИКАТ СООТВЕТСТВИЯ АУДИТОРА

Регистрационный номер № 04ЕАС1.СМ.03842-02

### НАСТОЯЩИЙ СЕРТИФИКАТ УДОСТОВЕРЯЕТ, ЧТО

### Гладун Виталий Викторович

требованиям добровольной сертификации «EAC соответствует системы предъявляемым к аудиторам внутренних проверок системы менеджмента качества изделий медицинских на соответствие стандарту ГОСТ ISO 13485-2017 (ISO 13485:2016) «Изделия Системные требования Системы менеджмента качества. для медицинские. регулирования»

Дата регистрации: 08-09-2021

Срок действия до: 07-09-2024

TO SPOBOLISHO

Руководитель органа по сертификации:

В. И. Погодин

Председатель

экспертной комисс

НАСТОЯЩИЙ СЕРТИФИКАТ ОБЯЗЫВАЕТ ОРГАНИЗАЦИЮ ПОДДЕРЖИВАТЬ СОСТОЯНИЕ ВЫПОЛНЯЕМЫХ РАБОТ В СООТВЕТСТВИИ С ВЫШЕУКАЗАННЫМИ СТАНДАРТАМИ, ЧТО БУДЕТ НАХОДИТЬСЯ ПОД КОНТРОЛЕМ ОРГАНА ПО СЕРТИФИКАЦИИ СИСТЕМЫ ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ "EAC AUDIT" И ПОДТВЕРЖДАТЬСЯ ПРИ ПРОХОЖДЕНИИ ЕЖЕГОДНОГО ИНСПЕКЦИОННОГО КОНТРОЛЯ



ФЕДЕРАЛЬНОЕ АГЕНТСТВО ПО ТЕХНИЧЕСКОМУ РЕГУЛИРОВАНИЮ И МЕТРОЛОГИИ

СИСТЕМА ДОБРОВОЛЬНОЙ СЕРТИФИКАЦИИ ГОСТ Р «EAC AUDIT»

РЕГИСТРАЦИОННЫЙ HOMEP POCC RU.32028.04EAC1

ОРГАН ПО СЕРТИФИКАЦИИ ООО «ГОРТЕСТ» РЕГИСТРАЦИОННЫЙ HOMEP POCC RU.32028

ИНН 7717616798 ОГРН 1087746489060

Юридический адрес: 109028, Россия, г. Москва, Серебряническая набережная, д. 27,

этаж 4, пом. 1, ком. 17

Телефон: 8 (800) 1000-730, e-mail: info@eacaudit.ru



### СЕРТИФИКАТ СООТВЕТСТВИЯ АУДИТОРА

Регистрационный номер № 04EAC1.СМ.03842-03

### НАСТОЯЩИЙ СЕРТИФИКАТ УДОСТОВЕРЯЕТ, ЧТО

### Нефуков Юрий Николаевич

сертификации «EAC добровольной требованиям соответствует системы предъявляемым к аудиторам внутренних проверок системы менеджмента качества изделий медицинских на соответствие стандарту ГОСТ ISO 13485-2017 (ISO 13485:2016) «Изделия Системные требования Системы менеджмента качества. медицинские. регулирования»

Дата регистрации: 08-09-2021

Срок действия до: 07-09-2024

Руководитель органа по сертификации:

В. И. Погодин

Председатель экспертной комиссии

POSPOBOJISHOV

Е. Д. Курбатова