



San Diego July 11th, 2018

We, ACON Laboratories Inc. having a registered office at 10125 Mesa Rim Road. San Diego, CA 92121, USA assign SRL Sanmedico having a registered office at A. Corobceanu street 7A, apt. 9, Chişinău MD-2012, Moldova , as authorized representative in correspondence with the conditions of directive 93/42/EEC, 98/79/EEC and 90/385/EEC.

We declare that the company mentioned above is authorized to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

ACON reserves the right to cancel this authorization at any time with a one month notice. If this is the case, ACON will honor any obligation to supply to our representative SanMedico SRL all the products distribution acquired or in the process of being acquired in Public Price bids and Public Tenders process.

Sincerely,


Jassy Alvarenga
Account Manager, International Sales



ACON Laboratories





Product Service

EC Certificate Full Quality Assurance System

Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6)
(List A and B and devices for self-testing)

No. V1 17 08 80997 017

Model(s): For Detail Models see attachment

Facility(ies): ACON Laboratories, Inc.
10125 Mesa Rim Road, San Diego CA 92121, USA

AZURE Institute, Inc.
10125 Mesa Rim Road, San Diego CA 92121, USA

ZERTIFIKAT ♦ CERTIFICATE ♦ CERTIFICADO ♦ CERTIFIKAT ♦ CEPTNΦNKAT ♦



Product Service

EC Certificate Full Quality Assurance System

Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6)
(List A and B and devices for self-testing)

No. V1 17 08 80997 017

Manufacturer: ACON Laboratories, Inc.
10125 Mesa Rim Road
San Diego CA 92121
USA



EC-Representative: Medical Device Safety Service GmbH

Schiffgraben 41
30175 Hannover
GERMANY

Product Category(ies):

In Vitro diagnostics for the detection of human infections and tumor markers, blood glucose measuring self-testing systems, self-testing devices for clinical chemistry, hematology and pregnancy

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device families in accordance with IVDD Annex IV. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of List A devices an additional Annex IV (4) certificate is mandatory. See also notes overleaf.

Report No.: SH17743EXT01

Valid from: 2017-09-13
Valid until: 2022-09-12



Date: 2017-08-30

S. Pinnit
Stefan Pinnit

TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

Page 1 of 4

TÜV SÜD Product Service GmbH, Zertifikatsstelle • Ritterstraße 65 • 80339 München | Germany

TÜV SÜD Product Service GmbH, Zertifikatsstelle • Ritterstraße 65 • 80339 München

Page 2 of 4



Storrum

ZERTIFIKAT ♦ CERTIFICATE ♦ CERTIFICADO ♦ CERTIFIKAT ♦ CEPTNΦNKAT ♦

For the product(s)/product category (ies):

- On Calli Plus Blood Glucose Monitoring System,
- On Calli Plus Blood Glucose Test Strips,
- On Calli EZ II Blood Glucose Monitoring System,
- On Calli Redi Blood Glucose Monitoring System,
- On Calli Redi II Blood Glucose Test Strips,
- On Calli Advanced Blood Glucose Monitoring System,
- On Calli Advanced Blood Glucose Test Strips,
- On Calli Platinum Blood Glucose Monitoring System,
- On Calli Platinum Blood Glucose Test Strips,
- On Calli Chosen Blood Glucose Monitoring System,
- On Calli Chosen Blood Glucose Test Strips,
- On Calli Vivid Blood Glucose Monitoring System (OGM-101),
- On Calli Vivid Blood Glucose Test Strips (OGS-101),
- On Calli Vivid Pal Blood Glucose Monitoring System (OGM-102),
- On Calli Sharp Blood Glucose Monitoring System (OGM-121),
- On Calli Sharp Blood Glucose Test Strips (OGS-121)
- On Calli Plus II Blood Glucose Monitoring System (OGM-171),
- On Calli Plus II Blood Glucose Test Strips (OGS-171),
- On Calli Extra Blood Glucose Monitoring System (OGM-191),
- On Calli Extra Blood Glucose Test Strips (OGS-191),
- On Calli GK Dual Blood Glucose & Ketone Monitoring System (OGM-161),
- On Calli Blood Ketone Test Strips (OGS-161),
- D-ONE Blood Glucose Monitoring System,
- D-ONE Blood Glucose Test Strips,
- Urnalysis Reagent Strips (Urine),
- UTI Urinary Tract Infection Test Strips
- Toxoplasma IgG EIA Test Kit,
- Toxoplasma IgM EIA Test Kit
- Rubella IgG EIA Test Kit,
- Rubella IgM EIA Test Kit,
- CMV IgG EIA Test Kit,
- CMV IgM EIA Test Kit,



- Total PSA EIA Test Kit,
- PT Coagulation Monitoring System (CCM-121),
- PT Coagulation Test Strips (CCS-121),
- Cholesterol Monitoring System (CCM-111),
- CHOL Total Cholesterol Test Devices (CCS-111),
- TRIG Triglycerides Test Devices (CCS-112),
- HDL High Density Lipoprotein Test Devices (CCS-113),
- 3-1 Lipid Panel Test Devices (CCS-114),
- Cholesterol CTRL Control Devices,
- Cholesterol Monitoring System (CCM-101),
- CHOL Total Cholesterol Test Strips (CCS-101),
- PT/INR Monitoring System (CCM-151),
- PT/INR Test Strips (CCS-151),
- Hemoglobin Testing System (CCM-141),
- Hemoglobin Test Strips (CCS-141),
- HCG Pregnancy Rapid Test Cassette (Urine),
- Pregnancy Rapid Test Midstream

Munich, MHS-CRT, 2017-08-30

S. Preis

Stefan Preis

Certification Medical Technology



S. Coornt

ZERTIFIKAT ◆ CERTIFICATE ◆ 認証証書 ◆ CERTIFICADO ◆ CERTIFICAT



Product Service

CERTIFICATE

No. Q1N 16 05 42074 027

Holder of Certificate: **Acon Biotech (Hangzhou) Co., Ltd.**
 No.210 Zhenzhong Road
 West Lake District
 310030 Hangzhou
 PEOPLE'S REPUBLIC OF CHINA

Facility(ies): Acon Biotech (Hangzhou) Co., Ltd.
 No.210 Zhenzhong Road, West Lake District,
 310030 Hangzhou, PEOPLE'S REPUBLIC OF CHINA



Certification Mark:



Scope of Certificate: **Design and Development,
 Production and Distribution of
 In Vitro Diagnostic Test Kits
 and Related Instruments,
 Lancet and Lancing Device**

Applied Standard(s): EN ISO 13485:2012 + AC:2012
 Medical devices - Quality management systems -
 Requirements for regulatory purposes
 (ISO 13485:2003 + Cor. 1:2009)
 DIN EN ISO 13485:2012

The Certification Body of TUV 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.: SH1610619
Valid from: 2016-07-15
Valid until: 2019-07-14



Date, 2016-07-08

 Stefan Preiß

Page 1 of 1

DAKKS
 Deutsche
 Institut für
 Zertifizierung

TUV SUD Product Service GmbH · Zertifizierstelle · Riederstraße 65 · 80339 München · Germany





LumiQuick Diagnostics, Inc.
2946 Scott Blvd., Santa Clara, CA 95054, USA

Tel: 1-408-855-0061
Fax: 1-408-855-0063
E-mail: info@lumiquick.com
Website: www.lumiquick.com

Date: February 13, 2018

LETTER OF AUTHORIZATION

To whom it may concern:

We, LumiQuick Diagnostics Inc. having a registered office at 2946 Scott Blvd, Santa Clara, CA 95054, USA, assign Sanmedico SRL having a registered office at str. A. Corobceanu 7A, apt. 9, Chişinău MD-2012, Moldova , as authorized representative in correspondence with the conditions of directive 98/79/EEC.

We declare that the company mentioned above is authorized to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

This authorization letter is valid until February 28, 2020.

Best regards,

Charles Yu
President





LumiQuick Diagnostics, Inc.
2946 Scott Blvd., Santa Clara, CA 95054, USA

Tel: 408-355-0064
Fax: 408-355-0064
E-mail: info@LumiQuick.com
Web: www.lumiquick.com

Declaration of Conformity

PRODUCT IDENTIFICATION		
Product name	Model/number	
Fecal Occult Blood Test Devices QuickProfile Fecal Occult Blood Test Card QuickProfile Fecal Occult Blood Test Strip	72001 72006	
MANUFACTURER		
Name of company	Address	Representative
LumiQuick Diagnostics, Inc.	2946 Scott Blvd. Santa Clara, CA 95054 USA	Jeff Wang
AUTHORIZED REPRESENTATIVE		
Name of company	Address	Telephone/email
Emergo Europe	Prinsessegracht 20 2514 AP The Hague, Netherlands	+31.70.345.8570 - phone +31.70.346.7299 - fax europe@emergogroup.com
CONFORMITY ASSESSMENT		
Device classification	Route to compliance	Standards applied
Class: Self-Certify	Annex III of IVDD 98/79/EC Council Directive	ISO 13485:2003

LumiQuick Diagnostics, Inc. declares that the above mentioned products meet the provision of the Council Directive 98/79/EC for In Vitro Diagnostic Medical Devices and Directive 98/79/EC as transposed in the national laws of the Member States.

COMPANY REPRESENTATIVE: Jeff Wang

TITLE: Quality Systems Manager

SIGNATURE: 

DATE: 28/04/2017

EC_Declaration_Letter_Emergo_E2R0_NewAddress



Handwritten signature



LumiQuick Diagnostics, Inc.
2946 Scott Blvd., Santa Clara, CA 95054, USA

Tel: 408-857-0001
Fax: 408-857-0863
E-mail: info@lumiquick.com
Web: www.lumiquick.com

Declaration of Conformity

PRODUCT IDENTIFICATION		
Product name	Model/number	
Cardiac Marker Test Devices		
QuickProfile Troponin I Serum Test Card	75001	
QuickProfile Troponin I Whole Blood Test Card	75002	
QuickProfile Cardiac Panel Serum Test Card	75003	
QuickProfile Cardiac Panel Whole Blood Test Card	75004	
QuickProfile Myoglobin Serum Test card	75005	
QuickProfile Myoglobin Whole Blood Test Card	75006	
QuickProfile CK-MB Serum Test Card	75007	
QuickProfile CK-MB Whole Blood Test Card	75008	
QuickProfile Troponin I Strip	75009	
QuickProfile CK-MB Strip	75010	
QuickProfile Myoglobin Strip	75011	
MANUFACTURER		
Name of company	Address	Representative
LumiQuick Diagnostics, Inc.	2946 Scott Blvd. Santa Clara, CA 95054 USA	Jeff Wang
AUTHORIZED REPRESENTATIVE		
Name of company	Address	Telephone/email
Emergo Europe	Prinsessegracht 20 2514 AP The Hague, Netherlands	+31.70.345.8570 - phone +31.70.346.7299 - fax europa@emergogroup.com
CONFORMITY ASSESSMENT		
Device classification	Route to compliance	Standards applied
Class: Self-Certify	Annex III of IVDD 98/79/EC Council Directive	ISO 13485:2003

LumiQuick Diagnostics, Inc. declares that the above mentioned products meet the provision of the Council Directive 98/79/EC for In Vitro Diagnostic Medical Devices and Directive 98/79/EC as transposed in the national laws of the Member States.

COMPANY REPRESENTATIVE: Jeff Wang

TITLE: Quality Systems Manager

DATE: 28/04/2017

SIGNATURE:

EC_Declaration_Letter_Emergo_E2R0_NewAddress





By Royal Charter

Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

LumiQuick Diagnostics, Inc.
2946 Scott Blvd
Santa Clara
California
95054
USA

Holds Certificate No:

FM 574919

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

The design, development, manufacture and distribution of in vitro diagnostics test kits and reagents used in the diagnosis and management of disease status, including Infectious Diseases tests, Drugs of Abuse tests, Cardiac Monitor tests, Cancer Marker tests, Fertility Hormone tests, ELISA tests & Urine Chemistry tests.

For and on behalf of BSI:

Stewart Brain, Head of Compliance & Risk - Medical Devices

Original Registration Date: 2011-10-20

Latest Revision Date: 2018-11-21

Effective Date: 2017-10-20

Expiry Date: 2020-10-19

Page: 1 of 1



...making excellence a habit.™

This certificate remains the property of BSI and shall be returned immediately upon request. An electronic certificate can be authenticated [online](http://www.bsigroup.com/ClientDirectory). Printed copies can be validated at www.bsigroup.com/ClientDirectory. To be read in conjunction with the scope above or the attached appendix.

Americas Headquarters: BSI Group America Inc., 12950 Worldgate Drive, Suite 800, Herndon, VA 20170-6007 USA
A Member of the BSI Group of Companies.



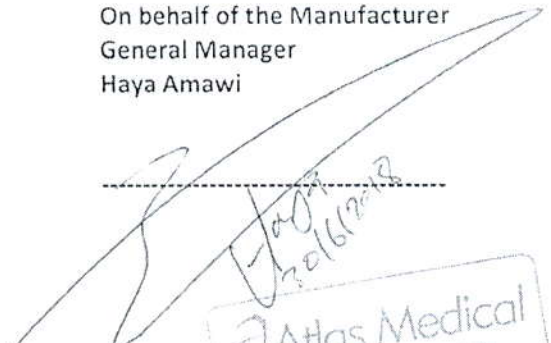
Date: 30/06/2018

STATEMENT

We, **Atlas Medical** having a registered office at William James House, Cowley Road, Cambridge, CB4 0WX, UK assign SRL Sanmedico having a registered office at A. Corobceanu street 7A, apt. 9, Chişinău MD-2012, Moldova, as authorized representative in correspondence with the conditions of directive 98/79/EEC.

We declare that the company mentioned above is authorized to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

On behalf of the Manufacturer
General Manager
Haya Amawi



Handwritten signature of Haya Amawi, dated 30/06/2018, over a dashed line. Below the signature is a rectangular stamp with the Atlas Medical logo and the text "Atlas Medical" and "Medical Products".

Head Office William James House, Cowley Rd, Cambridge, CB4 0WX, United Kingdom.
Tel: +44 (0) 1223 858910, Fax: +44 (0) 1223 858524

Middle East Site : King Abdullah the Second Industrial Estate, Street 19, Sahab Free Zone Area, P.O. Box: 204, Amman 11522, Jordan



CE Declaration of Conformity

According to Annex III of the IVD Directive 98/79/EC

We,

Atlas Medical

Head office: William James House, Cowley Road, Cambridge, CB4 0WX, UK

Tel: +44 1223 858 910

Fax: +44 1223 858 524

Email: info@atlas-site.co.uk

Middle East Site: Sahab Free Zone Area, P. O. Box 212555, Amman, Jordan.

Tel: +962 6 4026468

Fax: +962 6 4022588

Email: info@atlas-medical.com

Declare our responsibility that the following product:

See Attached list

- Comply with all essential requirements (Annex I) of the IVD Directive 98/79/EC. This compliance has been properly documented and covers the items listed in Annex I of the IVD Directive.
- This product is produced under Atlas quality system (ISO13485:2003) issued by Lloyd's Register Quality Assurance.
- Comply with the essential requirements of following standards (EN 18113-1, -2, -4:2011, EN ISO 15223:2012, EN ISO 13532, 2002, EN ISO 14971:2012, EN ISO 13640:2002, ISO 2859/1:1999, EN ISO 13612:2002, EN ISO 13641:2002).

And

Intended for In-Vitro Professional use only.

Manufacturer
Atlas Medical
 William James House, Cowley Rd.,
 Cambridge, CB4 0WX, UK

Catalogue No	Description	Catalogue No	Description
8.00.00	CRP latex Kits	8.02.48	Calcium Chloride
8.00.01	CRP latex Kits with buffer	8.02.69	Fibrinogen Reagent
8.00.02	ASO latex Kits	Hemoglobin Reagents	
8.00.03	ASO latex Kits with buffer	8.02.46	Drabkins Reagent, 40x
8.00.04	RF latex Kits	8.02.50	Hemoglobin Standard, 15g/dL
8.00.05	RF latex Kits with buffer	Sickle Cell Kits	
8.00.07	hCG Latex Kits	8.02.67	Sickle Cell Kit
8.00.08	IM (Hourse Stroma) Latex Kits	8.02.68	Sickle Cell positive & negative control set
8.00.11	SLE Latex kits	Urine Reagent Strips	
8.00.12	Staphylococcus Latex Kits	8.03.00	URS 1 Parameter: Glucose
8.00.13	Streptococcus Latex kits	8.03.01	URS 1 Parameter: Protein
8.00.15	E.Coli Latex Kits	8.03.02	URS 1 Parameter: Ketone
8.00.16	Rota Virus Latex Kits	8.03.03	URS 2 Parameters: Glucose, Ketone
8.00.17	D-Dimer Latex kits	8.03.04	URS 2 Parameters: Glucose, Protein
8.00.21	Waaler rose latex Kits	8.03.05	URS 2 Parameters: Urobilinogen, Bilirubin (Liver Function Test)
Febrile Antigen Kits			
8.01.00	Brucella Rose Bengal	8.03.06	URS 3 Parameters: Protein, pH, Glucose
8.01.01	Salmonella OA Reagent	8.03.07	URS 3 Parameters: Glucose, Protein, Ketone
8.01.02	Salmonella OB Reagent	8.03.15	URS 9 Parameters: Nitrite, Protein, pH, Blood, Specific Gravity, Ketone, Bilirubin, Glucose
8.01.03	Salmonella OC Reagent	8.03.16	URS 10 Parameters: Leukocytes, Nitrite, Urobilinogen, Protein, pH, Blood, Specific Gravity, Ketone, Bilirubin, Glucose
8.01.04	Salmonella OD Reagent	8.03.17	URS 10 Parameters: Nitrite, Urobilinogen, Protein, pH, Blood, Specific Gravity, Ketone, Bilirubin, Glucose, Ascorbic Acid
8.01.05	Salmonella HA Reagent	8.03.18	URS 11 Parameters: Leukocytes, Nitrite, Urobilinogen, Protein, pH, Blood, Specific Gravity, Ketone, Bilirubin, Glucose, Ascorbic Acid
8.01.06	Salmonella HB Reagent	Fertility Rapid Tests	
8.01.07	Salmonella HC Reagent	8.04.00	hCG Test Cassette, Urine
8.01.08	Salmonella HD Reagent	8.04.01	hCG Test Cassette, Urine/Serum
8.01.10	Brucella Abortus Reagent	8.04.04	hCG Test Strip, 5.0mm, Urine
8.01.11	Brucella Melitensis Reagent	8.04.05	hCG Test Strip, 3.5mm, Urine
8.01.12	Proteus OX2 Reagent	8.04.06	hCG Test Strip, 2.5mm, Urine
8.01.13	Proteus OX19 Reagent	8.04.10	hCG Test Strip, 5.0mm, Urine/Serum
8.01.14	Proteus OXK Reagent	8.04.12	hCG Test Strip, 3.5 mm, Urine/Serum
8.01.15	Brucella Antigen Kits	8.04.88	hCG Test Strip, 2.5 mm, Urine/Serum
8.01.16	Salmonella Antigen Sets	8.04.14	LH Test Cassette, Urine
8.01.17	Febrile Antigen Set (10 Antigens)	8.04.15	LH Test Strip, 3.5mm, Urine
8.01.17	Febrile Antigen Set (10 Antigens) With controls	Infectious Disease Rapid Test - Antibody Testing	
8.01.18	Salmonella Antigen Set, Widal Kit (6 Antigens: OA, OB, OD, HA, HB, HD)	8.04.20	H-pylori Antibody Test Cassette, Urynos Blood Serum/Plasma
8.01.18	Salmonella Antigen Set, Widal Kit (6 Antigens: OA, OB, OD, HA, HB, HD) with controls	CONTRACTE	
8.01.19	Febrile Antigen Positive Control	"SANMEDICO"	
8.01.20	Febrile Antigen Negative Control	S.R.L.	
Co-agglutination Reagents			
8.02.40	PT Calcium Rabbit Brain Thromboplastin, liquid	REPUBLICA ROMANA • YULIUS VERUS •	
8.02.41	APTT (PTT) Micronised Silica Platelet Substitute, Liquid	ID NO 100360307	
8.02.60	Normal Coagulation Control	SOCIETATEA CU RISP	
8.02.61	Abnormal Coagulation Control	1998	
8.02.44	PT Kit		
8.02.45	APTT (PTT) Kit		

Atlas Medical	Issue date	Date of review	Management approval	MRXDD10F.10
	December 2011	21st of March, 2018	<i>[Signature]</i>	08.02.2011



[Handwritten signature]

Certificate of Approval

This is to certify that the Management System of:

Atlas Medical

King Abdullah II Industrial Estate, Street No. 19, Sahab Free Zone Area, Amman, 11512, Jordan

has been approved by LRQA to the following standards:

ISO 13485:2003



Basem Obaid - Area Operations Manager

Issued By: Lloyd's Register EMEA

for and on behalf of: Lloyd's Register Quality Assurance Limited

Current Issue Date: 23 March 2018
Expiry Date: 31 March 2019
Certificate Issue Number: 10067833

Original Approvals:
ISO 13485 28 February 2009

Approval Certificate Number: ISO 13485 – 0046833

The scope of this approval is applicable to:
ISO 13485:2003

Design Manufacturing and Supply of Medical
Diagnostic Reagents and Kits



001



STATEMENT

We, "Technology-Standard" Ltd. having a registered office at 116/95, Kalinin Prospekt, Barnaul, 656037, Russia, assign SRL SANMEDICO having a registered office at A. Corobceanu street 7A, apt. 9, Chişinău MD-2012, Moldova, as authorized representative in correspondence with the conditions of directive 98/79/EC.

We declare that the company mentioned above is authorized to register, notify, renew or update the registration of medical devices on the territory of the Republic of Moldova.

"Tecnology-Standart" Ltd
116|95 Kalinin Prospekt
City of Barnaul, 656037, Russia
SRL SANMEDICO
A. Corobceanu street 7A, apt. 9,
Chişinău MD-2012, Moldova

Date: 01.12.2017

Director: Mr. A. B. *А. Б. Момот*

Signature: _____



ЗАЯВЛЕНИЕ

Мы, ООО «Технология-Стандарт», имеющее зарегистрированный офис по адресу 116/95, проспект Калинина, г. Барнаул, 656037, Россия, поручают SRL SANMEDICO, имеющую зарегистрированный офис на улице А.Коробчану 7А, кв. 9, Кишинёв MD-2012, Молдова, быть в качестве уполномоченного представителя в соответствии с условиями директивы 98/79/ЕС.

Мы заявляем, что упомянутая выше компания имеет право регистрировать, уведомлять, обновлять или возобновлять регистрацию медицинских изделий на территории Республики Молдова.

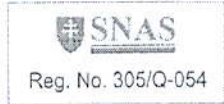
ООО Фирма «Технология-Стандарт»
656037 Россия г.Барнаул,
пр-кт Калинина 116/95
SRL SANMEDICO,
г. Кишинёв MD-2012, Молдова
ул. А.Коробчану 7А, кв. 9

Дата: 01.12.2017

Директор: А. Б. *А. Б. Момот*

Подпись: _____





CERTIFICATE

This certifies that the Quality management system for medical devices of company

«Technology-Standard» LTD

116/95, Kalinin Prospekt, City of Barnaul, 656037
RUSSIA

has been assessed by 3EC International and found to be in conformance with the following standard:

EN ISO 13485:2016

for the following scope:

DEVELOPMENT, PRODUCTION AND SALES OF DIAGNOSTIC KITS AND REAGENTS FOR IN VITRO DIAGNOSTICS OF HAEMOSTASIS SYSTEM

Certificate No.: M-0379/18

Date of issuance: July 26th, 2018

Original date of approval: August 5th, 2016

This certificate is valid from July 26th, 2018 to August 4th, 2019 on condition that organization will maintain effective Quality management system for medical devices. To verify the validity of this certificate please contact our office at: +421 (0)2 5831 8343.

This certificate fully supersedes previous certificate No. M-0379/16 issued on August 5th, 2016.

Issuing office: 3EC International a.s., Hraničná 18, 821 05 Bratislava, Slovak Republic

Dr. Katarina Tomin Srdošová
Head of Certification Body 3EC International a.s.

Certification body 3EC International a.s. is accredited by SNAS under registration number 305/Q-054 with accreditation certificate No. Q-054 for certification of Quality management systems for medical devices.



ЗАО «ЭКОлаб» 142530 Московская обл, г.Электрогорск, ул.Буденного, д.1
e-mail: sekretar@ekolab.ru, Сайт : www.ekolab.ru
Тел: (49643) 3-1374, 3-2311, факс (49643) 3-3143



ИНН: 5035025076, КПП: 503501001
Банк получателя: ПАО Сбербанк России г. Москва
в Орехово-Зуевском ОСБ № 1556/063
р/с 40702810040310124002
к/с 30101810400000000225
БИК 044525225

09.01.2019

АВТОРИЗАЦИЯ ДИСТРИБЬЮТОРА

Закрытое акционерное общество «ЭКОлаб» (Россия, 142530, Московская обл., г.Электрогорск, ул.Буденного, д.1) настоящим подтверждаем, что "SANMEDICO" SRL (ул. Коробчяну 7А, кв. 9, г. Кишинёв, Республика Молдова) является нашим эксклюзивным дистрибьютором и представителем в Республике Молдова и осуществляет участие с продукцией ЗАО «ЭКОлаб» в процедурах государственных закупок товаров на территории Республики Молдова, от своего имени ведет переговоры, представляет коммерческие предложения, заключает соответствующие договоры, а также осуществляет поставки указанной продукции на территории Республики Молдова.

Полномочия по настоящему авторизационному письму не могут быть переданы другим лицам.

Настоящее письмо действительно с момента подписания и до 31 декабря 2019г.

Генеральный директор



Борисов В.Ю.



Borisov



Certificat

Certificate

N° 2007/28641.5

AFNOR Certification certifies that the management system implemented by:
AFNOR Certification удостоверяет, что система менеджмента организации:



ЗАО «EKOlab»
ЗАО «ЭКОлаб»



for the following activities:
для следующих областей деятельности:

DEVELOPMENT, PRODUCTION, STORAGE AND SALE OF MEDICAL DEVICES
FOR IN-VITRO DIAGNOSTICS AND OF FINISHED MEDICINE

РАЗРАБОТКА, ПРОИЗВОДСТВО, ХРАНЕНИЕ И РЕАЛИЗАЦИЯ МЕДИЦИНСКИХ ИЗДЕЛИЙ ДЛЯ
IN-VITRO ДИАГНОСТИКИ И ЛЕКАРСТВЕННЫХ СРЕДСТВ

has been assessed and found to meet the requirements of:
проверена и признана соответствующей требованиям стандарта:

ISO 9001 : 2015

and is developed on the following locations:
и действует на следующих площадках:

142530, RUSSIA, MOSCOW REGION, ELEKTROGORSK CITY, Budennogo str., 1-1A
142530, РОССИЯ, МОСКОВСКАЯ ОБЛАСТЬ, г. ЭЛЕКТРОГОРСК, ул. Буденного, 1-1А

This certificate is valid from 2016-02-21.
Данный сертификат действителен с 2016-02-21 по:

2016-02-21

until
до

2019-02-21

Managing director of AFNOR Certification
Генеральный директор АФНОР Сертификейшн

F. LEBEUGLE



Certificat

Certificate

N° 2007/28642.4

AFNOR Certification certifies that the management system implemented by:
AFNOR Certification удостоверяет, что система менеджмента организации:



ЗАО «EKOlab»
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for the following activities:
для следующих областей деятельности:

DEVELOPMENT, PRODUCTION, STORAGE AND SALE OF MEDICAL DEVICES
FOR IN-VITRO DIAGNOSTICS.

РАЗРАБОТКА, ПРОИЗВОДСТВО, ХРАНЕНИЕ И РЕАЛИЗАЦИЯ МЕДИЦИНСКИХ ИЗДЕЛИЙ ДЛЯ
IN-VITRO ДИАГНОСТИКИ.

has been assessed and found to meet the requirements of:
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F. LEBEUGLE



afnor

11 rue de la République - 93011 La Plaine Saint-Denis Cedex - France - T: +33 (0)1 47 82 66 66 - F: +33 (0)1 47 82 66 20
Șosea de Republică nr.19, 06009 - București - România - T: +40 (0) 21 82 66 66 - F: +40 (0) 21 82 66 20