

ЗАО «ЭКОлаб» 142530 Московская обл, г.Электрoгорск, ул.Буденного, д.1  
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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

21.03.2018

### АВТОРИЗАЦИЯ ДИСТРИБЬЮТОРА

Закрытое акционерное общество «ЭКОлаб» (Россия, 142530, Московская обл., г.Электрoгорск, ул.Буденного, д.1) настоящим подтверждаем, что "SANMEDICO" SRL (ул. Коробчану 7А, кв. 9, г. Кишинёв, Республика Молдова) является нашим эксклюзивным дистрибьютором 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 2018г.

Генеральный директор



Борисов В.Ю.





# Certificat

Certificate

N° 2007/28641.5

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



**ZAO «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 площадях:

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

This certificate is valid from (date of issue):  
Данный сертификат действителен с (даты выдачи):

**2016-02-21**

and  
до

**2019-02-21**

Managing Director of AFNOR Certification  
Генеральный директор AFNOR Certification

**F. LEBEUGLE**

AFNOR Certification is a member of the AFNOR Group. AFNOR Certification is a member of the AFNOR Group. AFNOR Certification is a member of the AFNOR Group.



# Certificat

Certificate

N° 2007/28642.4

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



**ZAO «EKOlab»**  
ЗАО «ЭКОлаб»

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:  
проверена и признана соответствующей требованиям стандарта:

**ISO 13485 : 2003**

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

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

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Генеральный директор AFNOR Certification

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**EKOLab**

Declaration of Conformity

STED130-2017 vs. 01

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**DECLARATION OF CONFORMITY**

- 1) **Manufacturer** (Name, department): **CJSC EKOLab**  
Address: 1 Budennogo Str., Elektrogorsk, Moscow region, 142530, Russia
- 2) **European authorized representative: CEpartner4U BV**,  
Address: **ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS**,  
(on product labels printed as:  
CEpartner4U, ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS, www.cepartner4u.com)

- 3) **Product(s)** (name, type or model/batch number, etc.):

- Rabbit plasma

- 4) **The product(s) described above is in conformity with:**

Title	Document No.
In vitro Diagnostic Medical Devices Directive	98/79/EC

- 5) **Additional information** (conformity procedure, Notified Body, CE certificate, etc.):  
Conformity assessment procedure for CE marking: *In vitro* Diagnostic Medical Device Directive, Annex III  
Registration nr.: pending

Elektrogorsk, Russia; 2017-11-03

(Place & date of issue (yyyy-mm-dd))



V.Y. Borisov, General Director, CJSC EKOLab  
(name, function and signature of manufacturer)



**EKOLab**

Declaration of Conformity

STED130-2017 vs. 01

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**Appendix**

Date: 2017-11-08

List of devices:

Device name	Type/ model/ref number	Risk class / rule <sup>1</sup>	Code: EMDS/GMDN	First date of CE- compliance
Rabbit plasma		Low risk	15011290/0	2017-11-08

<sup>1</sup> See EDMS codes: <http://www.edms-ivd.be/> (products classification)/Preference GMDN





San Diego July 11<sup>th</sup>, 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



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ACON Laboratories





Product Service

# EC Certificate

## Full Quality Assurance System

Directive 93/79/EEC 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. Paus*

Shulan Paus

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

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TÜV SÜD Product Service GmbH · Zertifizierungsstelle · Riederstraße 65 · 80328 München · Germany

TÜV<sup>®</sup>

ZERTIFIKAT

CERTIFICADO

CEПTИΦИКАТ



CERTIFICATE

ZERTIFIKAT

11.08.17



Product Service

# EC Certificate

## Full Quality Assurance System

Directive 93/79/EEC 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



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TÜV SÜD Product Service GmbH · Zertifizierungsstelle · Riederstraße 65 · 80328 München



Product Service

Attachment for Certificate No V1 17 08 80997 017  
Supplement 001 dated 2017-08-30

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

- On Call Plus Blood Glucose Monitoring System,
- On Call Plus Blood Glucose Test Strips,
- On Call EZ II Blood Glucose Monitoring System,
- On Call Redi Blood Glucose Monitoring System,
- On Call Redi II Blood Glucose Test Strips,
- On Call Advanced Blood Glucose Monitoring System,
- On Call Advanced Blood Glucose Test Strips,
- On Call Platinum Blood Glucose Monitoring System,
- On Call Platinum Blood Glucose Test Strips,
- On Call Chosen Blood Glucose Monitoring System,
- On Call Chosen Blood Glucose Test Strips,
- On Call Vivid Blood Glucose Monitoring System (OGM-101),
- On Call Vivid Blood Glucose Test Strips (OGS-101),
- On Call Vivid Pet Blood Glucose Monitoring System (OGM-102),
- On Call Sharp Blood Glucose Monitoring System (OGM-121),
- On Call Sharp Blood Glucose Test Strips (OGS-121)
- On Call Plus II Blood Glucose Monitoring System (OGM-171),
- On Call Plus II Blood Glucose Test Strips (OGS-171),
- On Call Extra Blood Glucose Monitoring System (OGM-191),
- On Call Extra Blood Glucose Test Strips (OGS-191),
- On Call GK Dual Blood Glucose & Ketone Monitoring System (OGM-161),
- On Call Blood Ketone Test Strips (OGS-161),
- D-ONE Blood Glucose Monitoring System,
- D-ONE Blood Glucose Test Strips,
- Urinalysis 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.



Attachment for Certificate No V1 17 08 80997 017  
Supplement 001 dated 2017-08-30

**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

*I. P. P. P.*

Stefan Preiß

Certification Medical Technology



ZERTIFIKAT ◆ CERTIFICATE ◆ 認證證書 ◆ CERTIFICADO ◆ CERTIFICAT ◆ 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 TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). See also notes overleaf.

**Report No.:** SH1610619  
**Valid from:** 2016-07-15  
**Valid until:** 2019-07-14



**Date,** 2016-07-08  
  
Stefan Preiß

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## Declaration of Conformity

ACON Laboratories, Incorporated  
10125 Mesa Rim Road  
San Diego, CA 92121 USA

We declare under our sole responsibility that the *in vitro* diagnostic device:

Foresight HSV 1 IgG EIA Test Kit  
Foresight HSV 2 IgG EIA Test Kit  
Foresight HSV 1/2 IgG EIA Test Kit  
Foresight HSV 1 IgM EIA Test Kit  
Foresight HSV 2 IgM EIA Test Kit  
Foresight HSV 1/2 IgM EIA Test Kit

classified as others of the directive 98/79/EC,

meets all the provisions of the directive 98/79/EC on *in vitro* diagnostic medical devices which apply to it

This self-declaration is according to Annex III (excluding Section 6) of the Directive.

Authorized Representative:  
MDSS  
Schiffgraben 41  
30175 Hannover, Germany

Signed this 8<sup>th</sup> day of Oct, 2013  
in San Diego, CA USA



Qiyi Xie, MD, MPH  
Senior Staff, Regulatory Affairs  
ACON Laboratories, Inc.



10125 Mesa Rim Road · San Diego, CA 92121 · USA · Tel: (858) 875-8000 · Fax: (858) 875-8099  
E-mail: info@aconlabs.com







# XEMA

XEMA CO., Ltd.  
bdg. 48/4, 9th Parkovaya str., 105264, Moscow, Russia  
Tel./Fax: +7 (495) 510-57-07, +7 (495) 737-39-36  
E-mail: info@xema.ru, info@xema-medica.com  
Internet: www.xema.ru, www.xema-medica.com

## STATEMENT

We, XEMA Co., Ltd. having a registered office at 48, 9<sup>th</sup> Parkovaya st, 104264 Moscow, Russia, 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 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.

Date : November, 29, 2017

Signature:



Andrei P. Redkin

Deputy general manager



# Certificate

## Of Marketing Authorization of Medical Product

Nr. AR/IVMD/Xema/12-2016

Based on the basis of the Declaration of conformity and registration taking into account Article 10 of Directive 98/79/EC on In Vitro Diagnostic Medical Devices and Medical Devices Act (MPG) §§ 5,25,29,30

Ausgewählt auf Grund der Konformitätsklärung und Registrierung unter Berücksichtigung der Richtlinie 98/79/EG über In-vitro-Diagnostika und Medizinprodukte-gesetz (MPG) §§ 5,25,29,30

**Manufacturer:**  
Hersteller  
**Xema Co., Ltd.**  
bid.4, 48, The 8th Parkovskaya str.  
Moscow 105264, RUSSIA.  
info@zema.ru; www.zema.ru

**Product name:**  
Produkt  
Siehe Anhang zum Zertifikat

**Product Classification:**  
Produktklassifizierung  
In Vitro Diagnostic Medical Devices  
In-vitro-Diagnostika (IVD) Medizinprodukte

**Category:**  
Kategorie  
Common/ Other IVD  
Sonstige IVD-Produkte

**Conformity Module:**  
Konformitätsmodul  
Module A (EC Declaration of Conformity)  
(Annex III, except points, Directive 98/79/EC)  
Modul A (EG-Konformitätsklärung)  
(Anhang III, außer Nummer 6, Richtlinie 98/79/EG)

**Lead Competent Authority:**  
Zuständige Behörde  
DIMDI – German Institute of Medicinal Documentation and Information

**Product Registration Ref. No.:**  
(Per Article 10, Directive 98/79/EC)  
Produkt Registrierungsnummer  
(Gemäß Artikel 10 der Richtlinie 98/79/EG)

**Date of issue:** 2016-12-31  
Das Ausstellungsdatum

**Valid to:** 2019-12-31  
Gültig bis

**Represented in the EC by Polmed.de**  
Stelmacher S. 73773 Alchwald, Germany  
email: info@polmed.de  
tel: +49 711 52653379



*[Signature]*  
Polmed.de

Valid with the Annex of the Declaration of conformity and registration taking into account Article 10 of Directive 98/79/EC on In Vitro Diagnostic Medical Devices and Medical Devices Act (MPG) §§ 5,25,29,30

# Annex to the Certificate No.:

Anhang zum Zertifikat Nr.:

AR/IVMD/Xema/12-2016

The following medical devices can be placed on the market in the Federal Republic of Germany, in the member states of the European Economic Community (EEC) and in the other contract states of the agreement about the European Economic Area.

Die folgenden Medizinprodukte in der Bundesrepublik Deutschland, in den Mitgliedsstaaten der Europäischen Wirtschaftsgemeinschaft (EWG) und in den Vertragsstaaten der EÜG in den Vorlehr gebösch werden dürfen.

Nomenklature term Nomenklaturbezeichnung	Catalog number Katalognummer	Name of device Produktbezeichnung	DIMDI Product registration number Registrierungsnummer
1. THYROD PERIODICALLY (INCL. MICROZONMAI) ANTIBODIES	K131	4TPO EA Cat. Nr. K131	DE/CA37/IVD/13/44
2. THYROGLOBULIN AUTOANTIBODIES	K132	4TG EA Cat. Nr. K132	DE/CA37/IVD/13/43
3. MPO AMCA	K133	3MPO EA Cat. Nr. K133	DE/CA37/IVD/13/42
4. TISSUE TRANSGLUTAMINASE ANTIBODIES	K160 K161	AmH-TTG IGG EA Cat. Nr. K160 AmH-TTG IGA EA Cat. Nr. K161	DE/CA37/IVD/13/41
5. GLIADIN ANTIBODIES	K180 K181 K182A K182G	Gladin IGG EA Cat. Nr. K180 Gladin IGA EA Cat. Nr. K181 Diaminlated Gladin IGA EA, Diaminlated Gladin IGG EA	DE/CA37/IVD/13/40
6. IMMUNOGLOBULIN – TOTAL	K200	Total IGG EA Cat. Nr. K200	DE/CA37/IVD/13/39
7. THYROD STIMULATING HORMONE	K201 K201A K202	TSH EA Cat. Nr. K201 TSH Plus EA Cat. Nr. K201A LH EA Cat. Nr. K202	DE/CA37/IVD/13/38
8. LUTEINISING HORMONE	K203	FSH EA Cat. Nr. K203	DE/CA37/IVD/13/37
9. FOLLICLE STIMULATING HORMONE	K204	GH EA Cat. Nr. K204	DE/CA37/IVD/13/36
10. HUMAN GROWTH HORMONE	K205	HCG EA Cat. Nr. K205	DE/CA37/IVD/13/35
11. HUMAN CHORIONIC GONADOTROPIN TOTAL	K206	Progesterone EA Cat. Nr. K206	DE/CA37/IVD/13/34
12. PROGESTERONE	K207	Progesterone EA Cat. Nr. K207	DE/CA37/IVD/13/33
13. ESTRADIOL	K208	Estriol EA Cat. Nr. K208	DE/CA37/IVD/13/32
14. TESTOSTERONE (WITH DEHYDRO AND FREE TESTOSTERONE)	K209	Testosterone EA Cat. Nr. K209	DE/CA37/IVD/13/31
15. CORTISOL	K210	Salivary Testosterone EA Cortisol EA Cat. Nr. K210	DE/CA37/IVD/13/30
16. TRIODOTHYRONINE	K211	T3 EA Cat. Nr. K211	DE/CA37/IVD/13/29
17. THYRONINE	K212	T4 EA Cat. Nr. K212	DE/CA37/IVD/13/28
18. FREE TRIOODOTHYRONINE	K213	Free T3 EA Cat. Nr. K213	DE/CA37/IVD/13/27
19. FREE THYRONINE	K214	Free T4 EA Cat. Nr. K214	DE/CA37/IVD/13/26
20. DEHYDRO-EPANDROSTERONE SULPHATE (INCL. DHEA)	K215	DHEA-S EA Cat. Nr. K215	DE/CA37/IVD/13/25
21. 17 OH PROGESTERONE	K217	17-OH Progesterone EA Cat. Nr. K217	DE/CA37/IVD/13/24
22. CANCER ANTIGEN 125	K222	CA 125 EA Cat. Nr. K222	DE/CA37/IVD/13/23
23. CANCER ANTIGEN 15-9	K223	CA 19-9 EA Cat. Nr. K223	DE/CA37/IVD/13/22
24. CARCINOEMBRYONIC ANTIGEN	K224	CEA EA Cat. Nr. K224	DE/CA37/IVD/13/20

The above-mentioned medical products are marked with the CE symbol.  
Die oben genannten medizinischen Produkte sind mit dem CE-Zeichen gekennzeichnet.

Valid with the Annex of the Declaration of conformity and registration taking into account Article 10 of Directive 98/79/EC on In Vitro Diagnostic Medical Devices and Medical Devices Act (MPG) §§ 5,25,29,30



**Annex to the Certificate No.:**  
Anhang zum Zertifikat Nr.:

**AR/IVMD/Xema/12-2016**

The following medical devices can be placed on the market in the Federal Republic of Germany, in the member states of the European Economic Community (EEC) and in the other contract states of the agreement about the European Economic Area.

Die folgenden Medizinprodukte in der Bundesrepublik Deutschland, in den Mitgliedsstaaten der Europäischen Wirtschaftsgemeinschaft (EG) und in den Vertragsstaaten der EG in den Verkehr gebracht werden dürfen.

Nomenklature term Nomenklaturebezeichnung	Catalog number Katalognummer	Name of device Produktbezeichnung	DIAMDI Product registration number Spezialnummer
26. ALPHAFETOPROTEIN	K225	A1P EIA Cat. Nr. K225	DE/CA37/IVD/13/118
27. CANCER ANTIGEN 15-1	K226	M12 EIA (L5.1) EIA Cat. Nr. K226	DE/CA37/IVD/13/118
28. OTHER CANCER ANTIGENS	K227 K228	MUC11 M22 EIA Cat. Nr. K227; MUC11 M22 EIA Cat. Nr. K228	DE/CA37/IVD/13/117
29. OTHER OTHER TUMOUR MARKERS	K232	Thyroglobulin EIA Cat. Nr. K232	DE/CA37/IVD/13/116
30. HUMAN CHORIONIC GONADOTROPIN (INCL. SULFATE)	K235	Free beta HCG EIA Cat. Nr. K235	DE/CA37/IVD/13/115
31. PREGNANCY ASSOCIATED PLASMA PROTEIN-A (DOWNS)	K238	PAPP-A EIA Cat. Nr. K238	DE/CA37/IVD/13/114
32. OTHER OTHER PLASMA PROTEINS	K240	Albumin EIA Cat. Nr. K240	DE/CA37/IVD/13/113
33. C-REACTIVE PROTEIN	K250	CRP EIA Cat. Nr. K250	DE/CA37/IVD/13/112
34. SEX HORMONE BINDING GLOBULIN	K258	SHBG EIA Cat. Nr. K258	DE/CA37/IVD/13/111
35. TRYPHOSIN (T+J)	K291	Trypsin EIA Cat. Nr. K291	DE/CA37/IVD/13/110
36. IMMUNOGLOBULIN G	K271	Total IgG EIA Cat. Nr. K271	DE/CA37/IVD/13/109
37. IMMUNOGLOBULIN G SURCLASS REAGENTS	K272 K274	IgG2 EIA Cat. Nr. K272; IgG4 EIA Cat. Nr. K274	DE/CA37/IVD/13/108
38. IMMUNOGLOBULIN A	K275	Total IgA EIA Cat. Nr. K275	DE/CA37/IVD/13/107
39. IMMUNOGLOBULIN M	K277 K213	Total IgM EIA Cat. Nr. K277 Avidin/AT Immunology control set Cat. Nr. K213	DE/CA37/IVD/13/106
40. RHEUMATOID/ANTIMYELIN CONTROLS	K214 K215	Azidothymidine/Immunoassay control set Cat. Nr. K214; Azidothymidine/Immunoassay control set Cat. Nr. K215	DE/CA37/IVD/13/105
41. HORMONE CONTROLS	K221	Hormone immunoassay control set Cat. Nr. K221	DE/CA37/IVD/13/104
42. TUMOUR MARKER CONTROLS	K222	Onco/Obst immunoassay control set Cat. Nr. K222	DE/CA37/IVD/13/103
43. CYT6 22-1	K236	CYFRA 22-1 EIA	DE/CA37/IVD/13/102
44. CANCER ANTIGEN 72-4	K244	CA 72-4 EIA	DE/CA37/IVD/13/101
45. NEONATAL THYROID STIMULATING HORMONE	K201N	TSH-Neo EIA	DE/CA37/IVD/13/100
46. ESTRIOL	K218	Free Estriol EIA	DE/CA37/IVD/13/099
47. IMMUNOGLOBULIN E - MONOCLONAL/RESULIT - MULTI AG	K200S	Specific IgE EIA	DE/CA37/IVD/13/098
48. PAPPA AND LAMBDA CHAIN	K279E K279L	Free kappa IgE light chain EIA Free lambda IgE light chain EIA	DE/CA37/IVD/13/097
49. TRYPHOSIN NEONATAL	K242	Neonatal tryp EIA Cat. Nr. K242	DE/CA37/IVD/13/096
50. NEURON SPECIFIC ENDOSE	K234	NSE EIA Cat. Nr. K234	DE/CA37/IVD/13/095

The above mentioned medical products are marked with the CE symbol.  
Die oben genannten medizinischen Produkte sind mit dem CE-Zeichen gekennzeichnet.

Valid with the Federal Republic of Germany and all other states of the European Economic Community (EEC) and in the other contract states of the agreement about the European Economic Area.  
Gilt nur mit der Bundesrepublik Deutschland und allen anderen Staaten der Europäischen Wirtschaftsgemeinschaft (EWG) und in den Vertragsstaaten der EWG.

**Annex to the Certificate No.:**  
Anhang zum Zertifikat Nr.:

**AR/IVMD/Xema/12-2016**

The following medical devices can be placed on the market in the Federal Republic of Germany, in the member states of the European Economic Community (EEC) and in the other contract states of the agreement about the European Economic Area.

Die folgenden Medizinprodukte in der Bundesrepublik Deutschland, in den Mitgliedsstaaten der Europäischen Wirtschaftsgemeinschaft (EG) und in den Vertragsstaaten der EG in den Verkehr gebracht werden dürfen.

Nomenklature term Nomenklaturebezeichnung	Catalog number Katalognummer	Name of device Produktbezeichnung	DIAMDI Product registration number Spezialnummer
50. NEURON SPECIFIC ENDOSE	K234	NSE EIA Cat. Nr. K234	DE/CA37/IVD/13/095
51. OTHER OTHER TUMOUR MARKERS	K239	HE-4 EIA Cat. Nr. K239	DE/CA37/IVD/13/094
52. HIV IGG	K104	HIV 1/2 IGG EIA (Cat. Nr. K104)	DE/CA37/IVD/13/093
53. HIV IGM	K104M	HIV 1/2 IGM EIA (Cat. Nr. K104M)	DE/CA37/IVD/13/092
54. MYCOPLASMA ANTIBODY ASSAYS	K106	MycoSema IGG EIA (Cat. Nr. K106)	DE/CA37/IVD/13/091
55. SYPHILIS ANTIBODY ASSAYS TOTAL	K131	Treponema pallidum Total AB EIA (Cat. Nr. K131)	DE/CA37/IVD/13/090
56. SYPHILIS ANTIBODY IGG	K131G	Treponema pallidum IGG EIA (Cat. Nr. K131G)	DE/CA37/IVD/13/089
57. SYPHILIS ANTIBODY IGM	K131M	Treponema pallidum IGM EIA (Cat. Nr. K131M)	DE/CA37/IVD/13/088
58. H. PYLORI ANTIBODY ASSAYS	K139	h-pylori IGG EIA (Cat. Nr. K139)	DE/CA37/IVD/13/087
59. H. PYLORI ANTIBODY ASSAYS	K139M	h-pylori IGM EIA (Cat. Nr. K139M)	DE/CA37/IVD/13/086
60. ASPERGILLUS	K121	Aspergillus IGG EIA (Cat. Nr. K121)	DE/CA37/IVD/13/085
61. OTHER OTHER BACTERIOLOGY	K128	Ureaplasma IGG EIA (Cat. Nr. K128)	DE/CA37/IVD/13/084
62. GARDIA LAMBILIA	K171	Giardia lamblia Total AB EIA (Cat. Nr. K171)	DE/CA37/IVD/13/083
63. OTHER TUMOUR MARKER RAPID TESTS	K220V	XEMARapidScreen (Cat. Nr. K220V)	DE/CA37/IVD/13/082
64. OTHER TUMOUR MARKER RAPID TESTS	K222	XEMARapid25 (Cat. Nr. K222)	DE/CA37/IVD/13/081
65. OTHER TUMOUR MARKER RAPID TESTS	K239	XEMARapidE (Cat. Nr. K239)	DE/CA37/IVD/13/080
66. MAMMOGLOBULIN A IGA	K276	SECRETORY IGA (Cat. No. K276)	DE/CA37/IVD/13/079
67. LEPTOSPIROSIS	K175	Leptospira IGG EIA (Cat. No. K175)	DE/CA37/IVD/13/078
68. DISTOMATOSIS	K176	Parascia IGG EIA (Cat. No. K176)	DE/CA37/IVD/13/077
69. TESTOSTERONE (WITH DEHYDRO AND FREE TESTOSTERONE)	K219	Free Testosterone EIA (Cat. No. K219)	DE/CA37/IVD/13/076
70. HUMAN PLACENTAL LACTOGEN HPL	K246	Human Placental Lactogen DA (Cat. No. K246)	DE/CA37/IVD/13/075

The above mentioned medical products are marked with the CE symbol.  
Die oben genannten medizinischen Produkte sind mit dem CE-Zeichen gekennzeichnet.

Represented in the EC by Polmed.de  
Stelmecker S, 53775 Aichwald, Germany  
email: info@polmed.de  
tel: +49 711 52853279



Date: December 31, 2016



Valid with the Federal Republic of Germany and all other states of the European Economic Community (EEC) and in the other contract states of the agreement about the European Economic Area.  
Gilt nur mit der Bundesrepublik Deutschland und allen anderen Staaten der Europäischen Wirtschaftsgemeinschaft (EWG) und in den Vertragsstaaten der EWG.

# MANAGEMENT SYSTEM CERTIFICATE

Certificate No:  
53899-2009-AQ-MCW-FINAS

Initial certification date:  
21 May 2009

Valid:  
14 March 2018 - 28 February 2019

This is to certify that the management system of

## **XEMA CO., LTD.**

bldg. 48, 9-th Parkovaya str., Moscow, Russian Federation, 105264  
and the sites as mentioned in the appendix accompanying this certificate

has been found to conform to the Quality Management System standard:  
**ISO 13485:2003**

This certificate is valid for the following scope:  
**DESIGN, DEVELOPMENT, MANUFACTURING AND SALES OF KITS FOR IVD  
USE.**

Place and date:  
Moscow, 14 March 2018



For the issuing office:  
DNV GL - Business Assurance  
Trebkhprudny per. 9 build. 2, office 406,  
Moscow, Russian Federation

*S. Grov bme*  
Sergei Grov bme  
Management Representative

Lack of fulfilment of conditions set out in the Certification Agreement may render this Certificate invalid. This Certificate has been digitally signed. See [www.dnvgl.com/certificates](http://www.dnvgl.com/certificates) for more info.  
ACCREDITED UNIT: DNV GL BUSINESS ASSURANCE PJSC, ul. Muzhikova 3, 125080, Moscow, Russia. TEL: +7 495 10 33 4201. MAILBOX: DNV GL

Certificate No: 53899-2009-AQ-MCW-FINAS  
Place and date: Moscow, 14 March 2018

## Appendix to Certificate

### XEMA CO., LTD.

Locations included in the certification are as follows:

Site Name	Site Address	Site Scope
XEMA CO., LTD.	bldg. 48, 9-th Parkovaya str., Moscow, Russian Federation, 105264	DESIGN, DEVELOPMENT, MANUFACTURING AND SALES OF KITS FOR IVD USE.
XEMA Co., LTD ( production site)	Trubetskaya str., 2B, Balashkha, Moscow region, Russian Federation, 125000	DESIGN, DEVELOPMENT, MANUFACTURING AND SALES OF KITS FOR IVD USE.

Lack of fulfilment of conditions set out in the Certification Agreement may render this Certificate invalid. This Certificate has been digitally signed. See [www.dnvgl.com/certificates](http://www.dnvgl.com/certificates) for more info.  
ACCREDITED UNIT: DNV GL BUSINESS ASSURANCE PJSC, ul. Muzhikova 3, 125080, Moscow, Russia. TEL: +7 495 10 33 4201. MAILBOX: DNV GL



*Cleared*

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.

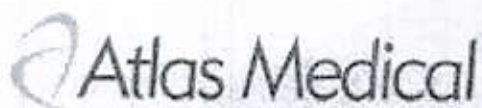
On behalf of the Manufacturer  
General Manager  
Haya Amawi



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 11552, Jordan





Declaration Ref No: DC11-0011

CE Declaration of Conformity

We,  
Atlas Medical

Head office: William James House, Cowley Road, Cambridge, CB0 4WX, UK  
Tel: +910 858 1223 44  
Fax: +524 858 1223 44  
Email: [info@atlas-site.co.uk](mailto:info@atlas-site.co.uk)

Middle East Site: Sahab Free Zone Area, P. O. Box 212555, Amman, Jordan.  
Tel.: +4026468 6 962  
Fax: +4022588 6 962  
Email: [info@atlas-medical.com](mailto:info@atlas-medical.com)

Declare our responsibility that the following product:

RPR Carbon Antigen

Is produced under Atlas quality system (ISO9001: 2008) and (ISO13485: 2003) supported by Lloyd's certificate and complies with the essential requirements of

In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex I  
And  
EN 18113-1, -2 :2011, EN ISO 15223:2012  
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.

This Declaration includes the batches produced beyond this day according to the product Lot Log.

Manufacturer  
Atlas Medical  
William James House, Cowley Rd.  
Cambridge, CB0 4WX, UK



Atlas Medical	First issue date	Date of review	Management approval	MRKDOT06
	Augsut-2003	06.11.2016	<i>S. Gull</i>	10 08.02.2013

# 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





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





# bsi.



By Royal Charter

## Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2003

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 13485:2003 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:

  
\_\_\_\_\_  
Carlos Pitanga, SVP, System Certification and Compliance

Original Registration Date: 2011-10-20

Latest Revision Date: 2017-10-09

Effective Date: 2017-10-20

Expiry Date: 2019-02-28

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](#). Printed copies can be validated at [www.bsigroup.com/ClientDirectory](http://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: 01<sup>st</sup> December 2017**STATEMENT**

We, **HiMedia Laboratories Pvt. Ltd.**, having a registered office at A-516, Swastik Disha Business Park, Via Vadhani Industrial Estate, L.B.S. Marg, Mumbai – 400 086, INDIA, 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.

**For HIMEDIA LABORATORIES PVT. LTD.,**



**Mr. V.M. SWARKE.**



**DIRECTOR – SALES & MARKETING**



**DECLARATION OF CONFORMITY**

1) Manufacturer (Name, department): HIMEDIA LABORATORIES PVT. LTD.  
Address: 23 Vadhani Industrial Estate, LBS Marg, Mumbai - 40, MS, India and

2) European authorized representative: CEpartner4U BV,  
Address: ELDOROLAAN 13, 3951DB MAAR, THE NETHERLANDS.  
CEpartner4U, ELDOROLAAN 13, 3951DB MAAR, THE NETHERLANDS. www.cepartner4u.eu

3) Product(s) (groupnames /):

Group	Group name	NL registration no.	No.
DCM&S	Dehydrated Culture Media & Supplements	NL-CA002-2013-26442	1
RPM	Ready Prepared Media Subgroups: Ready Prepared Plates, Ready Prepared Liquid & Solid Medium, Ready Prepared Slants, Ready Prepared Dual Media, HiDip Slides, HiGels Blood Culturing System, Transport Medium w/ swabs, Viral Transport Medium w/ swabs, L.J Medium Slants & Kits, Biochemical Kits for Mycobacteria, UTI Diagnostic Kits, Biochemical Identification Kits	NL-CA002-2013-26448	2
ESK	Epidemiological Screening Kit Subgroups: Hi Aureus Confirmation Kits	NL-CA002-2012-24117	3
ASS	Antimicrobial Susceptibility Systems Subgroups: Sensitivity Discs-Single & Multi Disc MIC Strips: HiComb Strips & Easy MIC Strips	NL-CA002-2013-26444	4
BDA	Bacteriological Differentiation Aids Subgroups: Ready-made Stains, Indicators & Reagents in liquid, Differentiation Discs & Strips, HiDried Rapid Identification Discs	NL-CA002-2013-26445	5
CCM	Cell Culture Media Subgroups: Karyotyping Media, Stem Cell Differentiation Media & Supplements, Stem Cell Freezing Medium, Stem Cell Differentiation Kits, Viral Transport Medium, Balanced Salt Solutions, Antibiotic solutions, Aerial Cell Culture Medium, Liquid	NL-CA002-2013-26446	6
MBP	Molecular Biology Products Subgroups: DNA & RNA Isolation Kits, Latex Agglutination Kits, Haematology Kits, Density gradient Separation Medium, PCR Kits	NL-CA002-2013-26447	7

type and model numbers: see appendix

4) The product(s) described above is in conformity with:

Title	Document No.
In vitro Diagnostic Medical Devices Directive	90/79/EEC

5) Additional information: (Conformity procedure, Modified Body, CE certificate, Registration nr., etc.):  
Conformity assessment procedure for CE marking: In vitro Diagnostic Medical Device Directive, Annex III

Mumbai, India, 2018-07-04

Dr. G.M. Wadnis, Managing Director  
(Name; function and signature of manufacturer)

(Place & date of issue (yyyy-mm-dd))

Declaration form: Standard ISO/IEC 17050-1:2010

DCM&S	FD003	Grift Microbacterial Supplement	Low risk	20/12/2012
DCM&S	FD004	GBS Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD005	G. Vaginalis Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD007	TTC Solution 1% (10 ml per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD009	Baitz Fuchsin (5.0 gm per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD009A	Litiera Selective Supplement (PALGAM) <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD062	Bacteroides Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD063	Litiera Selective Supplement II <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD063I	Litiera Selective Supplement II <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD066	Laptosira Enrichment <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD066	Sulpha Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD069	B.P Sulpha Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD070	McBride Litiera Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD071	Oxford Litiera Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD072	K1 Virulence Enrichment (20 ml per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD073	Diphtheria Virulence Supplement (Part A & B) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD075	Mycoplama Enrichment Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD082	Ampicilin Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD090	Campylobacter Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD091	Bruno Thymol Blue Supplement (20 mg per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD094	Trichomonas Selective Supplement II <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD095	10% Lactic Acid Solution (10 ml per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD099	Trichomonas Selective Supplement I <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD100	Muster Teichite Serum (5 ml per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD102	Tetracilin Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD103	Potassium Chlorate Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD105	Park and Sanders Selective Supplement II <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD106	Campylobacter Supplement VI (Butler) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD111	Enning Selective Supplement (Two Pack) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD112	George Enning Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD114	Vitamin K3 Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD117	Haemophilus Growth Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD118	Mucosal <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD119	Streptococci Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD120	Chlorotetracycline Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD126	Litiera Microbactan Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD130	Nadistic Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD135	CCDA Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD136	Litiera UVAM Supplement I <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD137	Litiera UVAM Supplement II <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD143	Legionella (DPIC) Selective Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD147	Tellurite - Culture Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD149	Neomycin Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD150	NYC Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD152	X174 Supplement <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD157	Urea SN (5 ml per vial) <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD158	Campylobacter Selective Supplement IV (Preston), Modified <td>Low risk <td>20/12/2012</td> </td>	Low risk <td>20/12/2012</td>	20/12/2012
DCM&S	FD159	Dorje's Antibiotic Supplement, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD160	Legionella (DPAN) Selective Supplement, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD161	Brucella Selective Supplement, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD163	Litiera Selective Supplement II, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD163-I	Litiera Selective Supplement II, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD164	Park and Sanders Selective Supplement I, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD165	Campylobacter Supplement - II (Butler), Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD169	CC Supplement, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018
DCM&S	FD171	McBride Litiera Supplement, Modified <td>Low risk <td>04/07/2018</td> </td>	Low risk <td>04/07/2018</td>	04/07/2018





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Nashik - 422 022, Maharashtra, India

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Date of initial issue: 29 December 2015

Valid until: 21 November 2020

Vienna, 22 November 2017

Quality Austria - Trainings, Zertifizierungs und Begutachtungs GmbH,  
AT-1010 Vienna, Zelinkagasse 10/3

*Scheiber*  
Konrad Scheiber  
General Manager

*Ar. Koubek*  
Dr. Mag. Anni Koubek  
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Registration No.: 00275/0

Date of initial issue: 21 November 2017

Valid until: 31 March 2019

Vienna, 22 November 2017

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AT-1010 Vienna, Zelinkagasse 103

*Konrad Scheiber*  
Konrad Scheiber  
General Manager

*Ing. Andreas Aichinger*  
Ing. Andreas Aichinger, MSc  
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 Awards this Certificate to

**HiMedia Laboratories Pvt. Ltd.**

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 Unit II : W-239(B), MIDC Phase II, Shivaji Udyog Nagar, Dombivli, District Thane - 421 204, Maharashtra, India  
 Unit III : D-61 MIDC, Phase-II, Near Shani Mandir, Dombivli, District Thane - 421204, Maharashtra, India



Unit I : Manufacture & supply of Biosciences products for applications in Microbiology (includes Dehydrated Culture Media, Culture Media Bases, Antimicrobial Susceptibility Systems & Bacteriological Differentiation Aids), Animal Cell Culture, Plant Tissue Culture and Molecular Biology Media  
 Unit II : Manufacture and supply of Sterile Ready Prepared Media  
 Unit III : Manufacture and supply of Sterile Ready Prepared Media

The validity of this Certificate will be maintained via annual surveillance audits and one renewal audit after three years.

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Report No.: QA/CA/WHO/069  
 Issue Date: 21/12/2016  
 Expiry Date: 20/12/2019  
 India - 20 Dec 2016  
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 Anil Kumar  
 Country Head

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