



STATEMENT

We, **Global Medikit Limited** having a registered office at 3.Dr.G.C Narang Marg, Delhi-110007, 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 93/42/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: 29.06.18

Signature: *R. Ravi*



Global Medikit Limited

Corporate Identity Number : U33119DL2001PLC110470

3, Dr. G. C. Narang Marg, Delhi - 110007 (INDIA) • Tel : +91-11-27662182 / 88 / 89 • Email : domestic@globalmedikit.in • Website : www.globalmedikit.in
Works : Khasra No. 323, (MI) Camp Road, Selaqui, Dehradun - 248197 (UTTRAKHAND)





EC DECLARATION OF CONFORMITY

According to section 2 of Annexure-V of the MDD 93/42/EEC concerning Medical Devices; we are;

Name of Manufacturer : Global Medikit Limited
Country of origin : India
Address / Tel / Fax : 3, Dr. G. C. Narang Marg, New Delhi, India -110007
Ph No. +91-11-27667888
Facility Address : Khasra No. 323 (MI), Central Hope Town, CAMP Road,
Selaqui Dehradun (Uttarakhand) India.

PRODUCT LIST

- Please refer to the attached Annex-A to Declaration of Conformity

Hereby declare under our own responsibility that the abovementioned products:

- Meet the provisions of the Council Directive 93/42/EC and the essential requirements which apply to them.
- The above mentioned devices have been classified as class III devices according to rule 6 & 7.
- This declaration is supported by the Quality Management System certification ISO 13485:2003 / NS - EN ISO 13485:2012, Certificate No. 248894 -2017-AQ-IND-NA-PS issued by DNV GL NEMKO PRESAFE AS - Veritasveien 3, N-1363 Hovik, Norway.
- This declaration is based on the conformity assessment of products to the requirements of Annex II according to the EC Conformity Certificate No. 247730 -2017- CE-IND-NA-PS issued by DNV GL NEMKO PRESAFE AS - Veritasveien 3, N-1363 Hovik, Norway.
- Notified Body No. : 2460.
- That the product comply all applicable harmonized standard ISO 9001:2008 / EN ISO 13485:2012; EN ISO 14971:2012; EN ISO 11135: 2014; EN ISO 10993-1: 2009, EN ISO 15223-1:2016, EN ISO 11737-1: 2006, EN ISO 11737-2:2009 and product Standard.
- The Product is manufactured according to Good Manufacturing Practices.

Issue place: Dehradun



Date: 30/11/2017

European Authorized Representative: Mr. Gideon ELKAYAM (CEO)

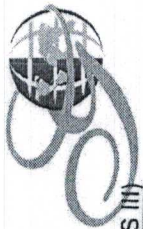
Registered Address:

Obelis s.a.
Bd. Général Wafis 53, B-1030 Brussels, Belgium
Phone: 32.2.732.59.54
Fax: 32.2.732.60.03
E-mail: mail@obelis.net

Valid until: This declaration is valid until further notice.

Global Medikit Limited

Corporate Identity Number: U33118DL2007PLC10470
3, Dr. G. C. Narang Marg, Delhi - 110007 (INDIA) • Tel : +91-11-27662182 / 88 / 89 • Email : domestic@globalmedikit.in • Website : www.globalmedikit.in
Works : Khasra No. 323, (MI) Camp Road, Selaqui, Dehradun - 248197 (UTTRAKHAND)



Annex A* – List of devices (CLASS III)

No.	Commercial name of Device	Generic Device Name and model	Class
1.	Epidural Mini Pack	Epidural Kit comprises Epidural catheter, LOR Syringe, Epidural Needle & Liquid Filter	III
2	Glospine	Spinal needle (Quincke Bevel & Pencil Point)	III
3	Glocont	Central Venous Catheter – Single Lumen (catheter through catheter technique)	III
4	Accont-1,2&3	Central venous catheter- Single, Double, Triple Lumen (Seldinger Technique)	III



Name of the Manufacturer

GLOBAL MEDIKIT LIMITED

Name and Position of the Signatory: Samar Keshari Jena (Assistant Manager QA)

Date: 30.11.2017

Stamp:



Global Medikit Limited

Corporate Identity Number: U33118DL2007PLC10470
3, Dr. G. C. Narang Marg, Delhi - 110007 (INDIA) • Tel : +91-11-27662182 / 88 / 89 • Email : domestic@globalmedikit.in • Website : www.globalmedikit.in
Works : Khasra No. 323, (MI) Camp Road, Selaqui, Dehradun - 248197 (UTTRAKHAND)



EC DECLARATION OF CONFORMITY

According to section 2 of Annexure-V of the MDD 93/42/EEC concerning Medical Devices, we are;

Name of Manufacturer : Global Medikit Limited
Country of origin : India
Address / Tel / Fax : 3, Dr. G. C. Narang Marg, New Delhi, India - 110007
Ph No. +91-11-27667888
Facility Address : Khasra No. 323 (MI), Central Hope Town, CAMP Road,
Selaqui Dehradun (Uttarakhand) India.

PRODUCT LIST

- Please refer to the attached Annex-A to Declaration of Conformity

Hereby declare under our own responsibility that the abovementioned products:

- Meet the provisions of the Council Directive 93/42/EC and the essential requirements which apply to them.
- The above mentioned devices have been classified as class Ila, I Ib & Is, devices according to rule 1, 2, 6 & 7.
- This declaration is supported by the Quality Management System certification ISO 13485:2003 / NS - EN ISO 13485:2012, Certificate No. 248894 -2017-AQ-IND-NA-PS issued by DNV GL NEMKO PRESAFE AS - Veritasveien 3, N-1363 Hovik, Norway.
- This declaration is based on the conformity assessment of products to the requirements of Annex II according to the EC Conformity Certificate No. 216144-2017- CE-IND-NA-PS issued by DNV GL NEMKO PRESAFE AS - Veritasveien 3, N-1363 Hovik, Norway.
- Notified Body No. : 2460.
- That the product comply all applicable harmonized standard ISO 9001:2008 / EN ISO 13485:2012; EN ISO 14971:2012; EN ISO 11135: 2014; EN ISO 10993-1:2009, EN ISO 15223-1:2016, EN ISO 11737-1: 2006, EN ISO 11737-2:2009 and product Standard .
- The Product is manufactured according to Good Manufacturing Practices.

Issue place: Dehradun



Samar Keshari Jena
[Assistant Manager QA]

Date: 30/11/2017

European Authorized Representative: Mr. Gideon ELKAYAM (CEO)

Registered Address:

Obelis s.a.
Bd. Général Watis 53, B-1030 Brussels, Belgium
Phone: 32.2.732.59.54
Fax: 32.2.732.60.03
E-mail: mail@obelis.net

Valid until: This declaration is valid until further notice.

Global Medikit Limited

Corporate Identity Number : U3319DL2007PLC10470
3, Dr. G. C. Narang Marg, Delhi - 110007 (INDIA) • Tel : +91-11-27662182 / 88 / 89 • Email : domestic@globalmedikit.in • Website : www.globalmedikit.in
Works : Khasra No. 323, (MI) Camp Road, Selaqui, Dehradun - 248197 (UTTARAKHAND)



Annex A* – List of devices(CLASS I, Ila&I Ib)

No.	Commercial name of Device	Generic Device Name and model	Class
1	Glomax	Tracheostomy Tube	I Ib
2	Heamodialysis Catheter	Heamodialysis Catheter Single, Double, Triple Lumen Central Venous Catheter (Seldinger technique)	I la
3	Glodrip	I. V. Infusion Set,	I la
4	Glodrip Alpha	I. V. Infusion Set with Airvent	I la
5	Glopreg	I. V Fluid Infusion Set with Flow regulator	I la
6	Glomic Alpha	Microdrip I. V infusion set with built in Airvent	I la
7	Gloflovi	I. V. Infusion Set with Airvent and Y- Site	I la
8	Gloryle	Ryle's Tube	I la
9	Glofant	Infant Feeding Tube	I la
10	Glonel	Nelaton Catheter	I la
11	Glolev	Stomach Tube	I la
12	Glosuc	Suction Catheter	I la
13	Glocase	Foley's Catheter	I la
14	Glokot	Malecot Catheter	I la
15	Gloyan	Yankauer handle with and without connecting tube	I la
16	Glofab	Laryngeal Mask Airway	I la
17	Gloflex	Three way Stop Cock	I la
18	Glomanifold	Stopcock manifold	I la
19	Gloflexo	Extension Lines with an Integrated three way stopcock	I la
20	Globob	Blood Administration Set	I la
21	Glovols	Measure Volume Burette Fluid Infusion Set	I la
22	Gloalpha	IV Catheter without Injection Valve & without wings	I la
23	Glocan	IV Catheter without Injection Valve & with wings	I la
24	Glofton	IV Catheter with Injection Valve & wings	I la
25	Glocan Alpha	IV Catheter without Injection Valve & without wings	I la
	Gloath	IV Catheter with Integrated 3-Way stopcock	I la

Global Medikit Limited



clacrum



27	Glonco	IV Catheter for Neonates	Ila
28	Glofancie	IV Catheter for Neonates	Ila
29	Gloveins	Extension Line with male and female ends	Ila
30	Gloveins Alpha	High Pressure Extension Line	Ila
31	Gloal	Endotracheal Tube (Plain)	Ila
32	Gloal Alpha	Endotracheal Tube (cuffed)	Ila
33	Glomask	Oxygen Mask (Adult, pediatric)	Ila
34	Glowin	Twin Bore Nasal Oxygen Set (Adult, Pediatric)	Ila
35	Gloreg	IV Flow Regulator	Ila
36	Gloex	Mucous Extractor	Ila
37	Glodrain	Wound Drainage Set	Ila
38	Glouro	Urine Bag	Is
39	Glometer	Urine Meter	Is
40	Glonutri	Enteral Feeding Bag	Is
41	Gloway	Guedel Airway (Oropharyngeal Airway)	Is
42	Luer Cap	Luer Cap	Is
43	Injection Stopper	Injection Stopper/NRV	Is
44	Glofin	A V Fistula Needle (With/Without Safety)	Ila
45	Glocic	Thoracic Drainage Catheter	Ila
46	Gloneb	Nebulizer Kit	Ila
47	Fabia	Percutaneous Sheath introducer Set	Ila

Name of the Manufacturer

GLOBAL MEDIKIT LIMITED

Name and Position of the Signatory: Samar Keshari Jena (Assistant Manager QA)

Date: 30.11.2017

Stamp:



Global Medikit Limited

Corporate Identity Number : U33130L2001PLC110470
3, Dr. G. C. Narang Marg, Delhi - 110007 (INDIA) • Tel : +91-11-27662182 / 88 / 89 • Email : domestic@globalmedikit.in • Website : www.globalmedikit.in
Works : Kharsa No. 323, (MI) Camp Road, Seisquit, Dohradun - 248197 (UTTARAKHAND)



EC Certificate Full Quality Assurance System

Certificate No.: 216144-2017-CE-IND-NA-PS Rev. 0.0 Project No.: PRJC-558191-2017-MSL-IND Valid Until: 09 October 2022

This is to certify that the quality system of:

Global Medikit Limited
Khasra No 323 (MI), Camp Road, Selaqui
248 197 Dehradun, Uttarakhand
India

For design, production and final product inspection/testing of:

Disposable Medical Devices

Has been assessed with respect to:

The conformity assessment procedure described in Article 11.3.a
and Annex II excluding section 4 (Module H2) of Council Directive
93/42/EEC on Medical Devices, as amended

and found to comply.

Further details of the product(s) and conditions for certification are given overleaf.

Place and Date:
Høvik, 09 October 2017

For:
DNV GL NEMKO PRESAFE AS



NOTIFIED BODY
PROD 021
Notified Body No.: 2460

Tone Kolpus

The Certificate has been digitally signed
See www.presafe.com/signatures for more info

Notice: The Certificate is subject to terms and conditions as set out in the Certification Agreement. Failure to comply may render this Certificate invalid.



EC Certificate Full Quality Assurance System

Certificate No.: 216144-2017-CE-IND-NA-PS Rev. 0.0 Project No.: PRJC-558191-2017-MSL-IND Valid Until: 09 October 2022

Jurisdiction

Application of Council Directive 93/42/EEC of 14 June 1993, adopted as "Forskrift om Medisinsk Utstyr" by the Norwegian Ministry of Health and Care Services.

Certificate history:

Revision	Description	Issue Date
0.0	Original Certificate	2017-10-09

Products covered by this Certificate:

Product Description	Product Name	Class
Tracheostomy Tube	Size: 2.0 to 11.0	I Ib
Haemodialysis Catheter	Single Double, Triple Lumen Central Venous Catheter (Seldinger technique)	I Ia
I.V. Sets	I.V. Infusion Set, I.V. Infusion Set with Airvent, Microdrip infusion set with built in Airvent, I.V. Set With Flow Regulator, I.V. Infusion Set with Airvent and Y-Site	I Ia
Ryle's Tube	Size(FG):8,10,12,14,16,18,20	I Ia
Infant Feeding Tube	Size(FG):4,5,6,8,10,12,14,16,18	I Ia
Nelaton Catheter	Size(FG):6,8,10,12,14,16,18,20	I Ia
Stomach Tube	Size(FG):8,10,12,14,16,18,20,22,24	I Ia
Suction Catheter	Size(FG):6,8,10,12,14,16,18,20,22,24	I Ia
Foley's Catheter	Latex Foley Catheter (Two Way/ Three Way) Silicone Foley Catheter (Two Way/ Three Way)	I Ia
Malecot Catheter	Size:8Fr to 42Fr	I Ia
Yankauer handle with and without connecting tube	Crown tip, Plain tip	I Ia
Laryngeal Mask Airway	Size: 1 to 5	I Ia
Stop Cock / Stopcock manifold	3-way stop Cock (Plain & Lipid Resistant) 3-way stop Cock (Plain & Lipid Resistant) With	I Ia





EC Certificate
Full Quality Assurance System

Certificate No.: 216144-2017-CE-IND-NA-PS Rev. 0.0
Project No.: PRJC-558191-2017-MSL-IND
Valid Until: 9 October 2022

Certificate No.: 216144-2017-CE-IND-NA-PS Rev. 0.0
Project No.: PRJC-558191-2017-MSL-IND
Valid Until: 9 October 2022

	extension tube, 2 Way,3 Way,4 Way, Stopcock in line (Plain & Lipid Resistant)		
Blood Administration Set	Vented /Non-Vented	Ila	
Measure Volume Burette	100ml, 110ml, 150ml	Ila	
Fluid Infusion Set	IV Catheter without injection valve & wings, IV Catheter without injection valve & with wings IV Catheter with injection valve & with wings IV Catheter with integrated 3 way stop cock IV Catheter for Infants and Neonates IV Catheter for Neonates Safety IV Catheter with injection valve and wings	Ila	
I.V. Catheter			
Extension Set/Tube	Extension Line With male and female ends (Single Lumen, Double Lumen, Triple Lumen), High Pressure Extension Line With male and female Ends	Ila	
Endotracheal Tube	Endotracheal Tube with Cuffed Endotracheal Tube without Cuffed	Ila	
Oxygen Mask & Twin Bore Set	Oxygen Mask with tubing and nose clip (Adult, Pediatric) Twin Bore Nasal Oxygen Set (Adult, Pediatric)	Ila	
IV Flow Regulator	5 – 300ml/h	Ila	
Mucous Extractor	8FG, 10FG, 12FG, 14FG,	Ila	
Wound Drainage Set	6 FG TO 18 FG	Ila	
Urine Bag	Adult, Paediatric, Legbag	Is	
Urine Meter	Urine Collection Bag With Measured Volume Meter	Is	
Enteral Feeding Bag		Is	
Guedel Airway (Oropharyngeal Airway)	Size: 000,00,0,1,1,5,2,3,4,5,6	Is	
Luer Cap		Is	

The complete list of devices is filed with the Notified Body

Sites covered by this certificate

Site Name	Address
Global Medikit Limited	Works: Khasra No.323 (MI), Camp Road, Selaqui, 248 197 Dehradun, Uttarakhand, India Regd Office: 3, Dr.G.C. Narang Marg, Delhi, 110007 India

EU Representative

Obelis s.a., Brussels, Belgium



Valid Until:
9 October 2022

The certificate is subject to the following terms and conditions:

- Any producer (see 2001/95/EC for a precise definition) is liable for damage caused by a defect in his product(s), in accordance with directive 85/374/EEC, as amended, concerning liability of defective products.
- The certificate is only valid for the products and/or manufacturing premises listed above.
- The Manufacturer shall fulfil the obligations arising out of the quality system as approved and uphold it so that it remains adequate and efficient.
- The Manufacturer shall inform Presafe of any intended updating of the quality system and Presafe will assess the changes and decide if the certificate remains valid.
- Periodical audits will be held, in order to verify that the Manufacturer maintains and applies the quality system. Presafe reserves the right, on a spot basis or based on suspicion, to pay unannounced visits.

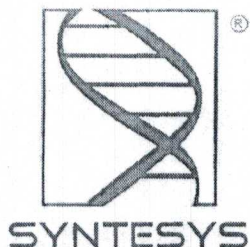
- Changes in the quality system affecting production.

- Periodical audits not held within the allowed time window.

When meeting with the terms and conditions above, the producer may draw up an EC declaration of conformity and legally affix the CE mark followed by the Notified Body identification number of Presafe.

MSD-CO-078 DNV GL NEMKO PRESAFE AS · Ventasveien 3, N-1363 Høvik, Norway · Registered Enterprise No: NO 997 067 401 MVA Page 5 of 5





SYNTESYS S.A.S. DI RINALDO R. & C.

VIA G. GALILEI, 10/3
35037 Z.I. SELVE DI TEOLO (PD)
TEL. +39 049 9903866 R.A. FAX +39 049 9903867
COD. FISCALE P.IVA N. REG. IMP. PADOVA 03573950288
E-MAIL INFO@SYNTESYS.IT - WEB WWW.SYNTESYS.IT

AUTHORIZATION LETTER

We, **Syntesys S.A.S.** having a registered office at Via G. Galilei 10/3, 35037 Selve di Teolo - PD - Italy, assign **Sanmedico SRL** having a registered office at A. Corobceanu str., apt. 9, Chişinău MD-2012, Moldova, as authorized representative in correspondence with the conditions of directive 98/79/CE and 93/42/CE.

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.

Teolo, 02.01.2018

 **SYNTESYS S.A.S.**
DI RINALDO R. & C.
Via G. GALILEI, 10/3
Z.I. SELVE 35037 TEOLO (PD) - CE 03573950288
TEL. 049/9903866 R.A. FAX 049/9903867

Rinaldo Ruggero
CEO and Legal Representative
SYNTESYS S.A.S.





SYNTEYS S.A.S. DIRINALDOR & C
6 GALLIEI, 10/3
35037 ZI. SELVE DI TEOLO (PD)
TEL. +39 049 9903866 R.A. FAX. +39 049 9903867
COD. FISCALE PIVA N. REG. IMP. PADOVA 03573950288
E-MAIL: INFO@SYNTEYS.IT - WEB: WWW.SYNTEYS.IT

SYNTESIS

produttore/manifatturere

SYNTESYS S.a.s. di Rinaldo Ruggero & C.
indirizzo/address

Via G. Galilei, 10/E 35037 Zona Industriale SELVE DI TEOLO (PADOVA) ITALY

0 rappresentante il mandatario autorizzato entro la Unione Europea o rappresenting the authorized mandatary within the European Community

Mandatario autorizzato/authorized mandatary

indirizzo address

Dichiara sotto la propria responsabilità che il prodotto/declares under his own responsibility that the product:

Denominazione degli

Articoli	Prodotti/Description of	Manufacturer
1	...	...
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Contenitori per urina, contenitori per feci, contenitori universali, Pipette Pasteur, Piastre di Petri, Anse Sterili per batteriologia, Aste a "L", Punta di Eppendorf gialli e blue, cuvette per spettrofotometro, tazzine per campionamento siero, siringhe a cassetta per distacco ed estrazione del coagulo, lancette in polistirolo monouso, provette monouso in polistirolo, tappi alettati per provette diam. 12 mm e 16 mm, provette con granuli ed acceleratore, provette sottovuoto per prelievo, Sistema SEDIPLAST, microprovette, portavetrini, vetrini precolorati, portaprovette, supporti per microprovette, bottiglie per raccolta urine.

Urine container, faeces container, universal container, Pasteur pipette, Petri dishes, Sterile loops, Sterile loops open "L", Eppendorf tips yellow and blue, cuvettes for spectrophotometer, samples cups, Rod to detach clot, disposable forceps, Disposable plastic tubes, winged stoppers for tubes diam. 12mm & 16mm, Test tube with granules and clot activator, vacuum test tube, VEDIPLAST system, micro test tubes, Slides Hailer, "TESTSIMPLETS" slide, Rack for test tubes, rack for micro test tubes, Bottles for urine collection.

Polipropilene, Polistirolo, Polietilene e
Polimetilmetacrilato

Material/Material

Polypropylene, Polystyrene, Polyethylene and
Polymethylmetacrylate

è conforme alle disposizioni della direttiva 98/79/CE concernente i dispositivi medici diagnostici in vitro e recepito in Italia con D.L. del 06/09/2000 n° 332 allegato I (requisiti essenziali) ed è fabbricato in accordo ai requisiti di cui all'Allegato III della sopra citata direttiva / It meets the CE Directive 98/79 CE about in vitro diagnostic device specifications established by the Italian law n° 332, dated 6th September 2000. The device is made according to the specifications of the III attached of the above-mentioned directive.

Dichiara inoltre che la documentazione tecnica di supporto alla presente dichiarazione di conformità è conservata presso gli uffici dell'azienda e sarà posta alla disposizione di chi la richiederà/declares that all technical documents attached to this conformity statement are filed in our company and can be consulted by any authorized body on demand.

9702
Issued on January 7th 2016
Data 07/01/2016

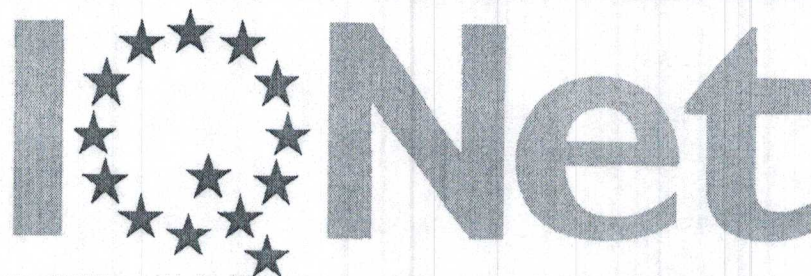
SYNTESIS S.A.S.

Il legale rappresentante
Rinaldo Ruggero



PRODOTTI E STRUMENTAZIONE PER ANALISI DI LABORATORIO • DISPOSABILE LABWARE

PRODOTTI E STRUMENTAZIONE PER ANALISI DI LABORATORIO - DISPOS



THE INTERNATIONAL CERTIFICATION NETWORK

CERTIFICATE

CISQ/ICIM SPA has issued an IQNet recognized certificate that the organization:

SYNTESYS S.a.s. di Rinaldo Ruggero e C.

Via G. Galilei, 10/3 - Zona Industriale - I-35037 Selve di Teolo (PD)

has implemented and maintains a

Quality Management System

for the following scope:

Trading of products for laboratory analysis. Design, manufacturing and sale of products for laboratory analysis and sanitary products. Sale agency of instruments, reagents and consumable products for laboratory diagnostic.

which fulfils the requirements of the following standard:

ISO 9001:2015

Issued on: **2018-06-04**

First issued on: **2013-06-05**

Expires on: **2019-06-04**

This attestation is directly linked to the IQNet Partner's original certificate and shall not be used as a stand-alone document.

Registration Number: **IT-83562**



Alex Stoichitoiu
President of IQNET



Ing. Claudio Provetti
President of CISQ

IQNet Partners*:

AENOR Spain AFNOR Certification France APCER Portugal CCC Cyprus CISQ Italy
CQC China CQM China CQS Czech Republic Cro Cert Croatia DQS Holding GmbH Germany FCAV Brazil
FONDONORMA Venezuela ICONTEC Colombia Inspecta Sertifiointi Oy Finland INTECO Costa Rica
IRAM Argentina JQA Japan KFQ Korea MIRTEC Greece MSZT Hungary Nemko AS Norway NSAI Ireland
NYCE-SIGE México PCBC Poland Quality Austria Austria RR Russia SII Israel SIQ Slovenia
SIRIM QAS International Malaysia SQS Switzerland SRAC Romania TEST St Petersburg Russia TSE Turkey YUQS S2
IQNet is represented in the USA by: AFNOR Certification, CISQ, DQS Holding GmbH and NSAI Inc.



* The list of IQNet partners is valid at the time of issue of this certificate. Updated information is available under www.iqnet-certification.com



THE INTERNATIONAL CERTIFICATION NETWORK

CERTIFICATE

CISQ/ICIM SPA has issued an IQNet recognized certificate that the organization:

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Via G. Galilei, 10/3 - Zona Industriale - I-35037 Selve di Teolo (PD)

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Quality Management System

for the following scope:

Trading of products for laboratory analysis. Design, manufacturing and sale of products for laboratory analysis and sanitary products. Sale agency of instruments, reagents and consumable products for laboratory diagnostic.

which fulfils the requirements of the following standard:

UNI CEI EN ISO 13485:2016

Issued on: **2018-06-04**

First issued on: **2014-06-21**

Expires on: **2019-06-04**

This attestation is directly linked to the IQNet Partner's original certificate and shall not be used as a stand-alone document.

Registration Number: **IT-93779**



Alex Stoichitoiu
President of IQNET



Ing. Claudio Provetti
President of CISQ

IQNet Partners*:

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