EC Certificate Full Quality Assurance System: Certificate GB19/964232



The management system of

Intersurgical Ltd.

Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, UK has been assessed and certified as meeting the requirements of

Directive 93/42/EEC

on medical devices, Annex II (excluding Section 4)

For the following products

The scope of registration appears on page 2 of this certificate.

This certificate is valid from 24 May 2021 until 26 November 2023 and remains valid subject to satisfactory surveillance audits.

Issue 8. Certified since 11 January 1995

Certification is based on reports numbered GB/PC 04303

This is a multi-site certification. Additional site details are listed on the subsequent page.

Authorised by

Global Medical Devices Head of Notified Body

SGS Belgium NV, Notified Body 1639

SGS House Noorderlaan 87 2030 Antwerp Belgium t +32 (0)3 545-48-48 f +32 (0)3 545-48-49 www.sgs.com

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Intersurgical Ltd. Directive 93/42/EEC

on medical devices, Annex II (excluding Section 4)

Issue 8

Detailed scope Sterile and Non-Sterile medical devices for respiratory care, in the areas of airway management, anaesthesia, critical care, oxygen and aerosol therapy: Sterile and Non-Sterile Anaesthetic Breathing Systems Aerosol and Oxygen Face Masks **Anaesthetic Face Masks** Sterile Endotracheal Tube Introducer and Sterile Airway Stylets Sterile and Non-Sterile Breathing Systems Non-Heated Respiratory Bubble Humidifier Sterile and Non-Sterile Catheter Mounts Sterile and Non-Sterile Breathing System Connectors Sterile and Non-Sterile Respiratory Filters Breathing System Flexible Tubing High Concentration Oxygen Face Masks Sterile and Non-Sterile Heat and Moisture Exchangers Sterile and Non-Sterile HME Filters Sterile and Non-Sterile Inspiratory Line Humidification Chambers Sterile I-gel Supraglottic Airways Sterile Laryngeal Airways Gas Sampling/Monitoring Respiratory Tubing Sterile and Non-Sterile Heated Wire Breathing Systems. Heated Wires and attachments (electrical adaptor leads) **Electrically Powered Moisture Condenser, Nasal Cannulae Nebulising System Delivery Sets** Suction and Irrigation Oral Care Toothbrush Oxygen Administration Tubing Repeated Use Breathing Systems **Breathing Systems Reservoir Bags** Manual Pulmonary Resuscitation Systems Carbon Dioxide Absorbents Sterile and Non-Sterile Tracheal Suction Systems Sterile Endotracheal Tubes Venturi Valves and Venturi Valve Face Mask Kits Wall Humidifier Nebuliser **Breathing System Water Traps CPAP Bi-level Nasal Masks and NIV Face Masks** Pressure Limiting Valves

Peep Valves One Way Directional Valves
Infant Nasal CPAP Breathing System
Oxygen Recovery Kits
Endoscopy Molar Bite Block
Carbon Dioxide Cuvette

Class I sterile: Sterility aspects only - Restricted to the aspects of manufacture concerned with securing and maintaining sterile conditions:

Sterile Guedel Airways

Certificate GB19/964232 continued





Intersurgical Ltd. Directive 93/42/EEC

on medical devices, Annex II (excluding Section 4)

Issue 8

Detailed scope

Additional facilities

Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, UK

Unit 3, Molly Millars Bridge, RG41 2WY, UK

Dray House, Molly Millars Lane, RG41 2PX, UK

Brook House, Molly Millars Bridge,, RG41 2WY, UK

Unit 1, Molly Millars Lane, RG41 2QZ, UK

Circuit House, Pitronnerie Road, Industrial Estate, St Peter Port, Guernsey, GY1 2RL, UK

UAB Intersurgical Arnionių g.60, LT-18170 Pabradė, Lithuania

Arnionių g. 60A, Pabradė, LT-18170, Lithuania

Arnionių g. 45, Pabradė, LT-18170, Lithuania

Duksto kelias 84A, Visaginas, LT-31146, Lithuania



Quality, innovation and choice

Intersurgical Ltd, Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, UK
T: +44 (0)118 9656 300 F: +44 (0)118 9656 356 info@intersurgical.com www.intersurgical.com

claration of	Conformity		

We, Intersurgical Ltd (address: Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, United Kingdom) as manufacturer, hereby declare under the sole responsibility of that the below mentioned devices comply with European Medical Devices Directive 93/42/EEC with all amendments and transposing legislation.

Authorised Representative in the European Economic Area (EEA): UAB Intersurgical (address: Arnioniu g. 60, Pabradė, LT-18170, Lithuania)

Carbon Dioxide Absorbents

These are class IIA medical devices, in accordance with rule 2 of Annex IX of the Medical Devices Directive 93/42/EEC (classification.doc)

GMDN codes - 42414, 36051

Essential requirements checklist is on IQR139.

Internally manufactured components master data is on EFACS (Parts Master) and design drawings are on IQR69 (Product and Mould Drawing List).

Internally manufactured finished products master data, production drawings and Master Product Formulae are on EFACS (Master Product Formula, Parts Master) and IQR69 (Product and Mould Drawing List).

Externally manufactured materials, components, design drawings and products are on EFACS (Parts Master), IQR98 (Incoming Product Specification) and IQR69 (Product and Mould Drawing List).

Instructions for use, pack inserts and labels are as detailed on EFACS (Master Product Formula) and IQR107 (Index of Controlled Artworks).

Label content is on EFACS (Master Product Formula) and Labels Printing System.

Product realisation processes are referenced in IQM section 7.1

Product Codes

As listed in EFACS MPF Details under group DCSODA.DOC.

This range is subject to the procedure set out in Annex 2 excluding section 4 of Directive 93/42/EEC as amended by 2007/47/EC, under the supervision of Notified Body Number 1639, SGS Belgium NV, SGS House Noorderlaan 87-2030, Antwerpen, Belgium

EC Certificate Full Quality Assurance System GB19/964232 has been issued for the management system of Intersurgical Ltd, which has been assessed and certified according to the requirements of Annex 2 excluding section 4 of Directive 93/42/EEC as amended by 2007/47/EC.

Ivan Seniut

Group Quality and Regulatory Affairs Director Duly authorised for and on behalf of Intersurgical Ltd Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, United Kingdom

Issue 20 Valid from 1 January 2021 DCSODA.DOC



Appendix for EU Declaration of Conformity Group: DCSODA.DOC

Product Codes:

	Vicinity of the second			
2169001	2169002	2169003	2172000	2173000
2174000	2175000	2178000	2179000	2180000
2186000	2187000	2188000	2191001	2192001
2193001	2196000	2197000	2198000	2199001
2199002	2199003	1855 (1917)	olt, spec 2 (20) to	(ac) let leon

in granded by 2007/47/17C, under the supervision of Various Body Marker 1679, SUS Display MV







Manufacturer's Authorization

Date: 01.08.2022

To Whom It May Concern

We Intersurgical Ltd, located at Crane House, Molly Millars Lane, Wokingham, Berkshire, RG41 2RZ, United Kingdom are the registered legal manufacturer of medical device for respiratory support. We have factories located at UAB Intersurgical, Arnoniu, g.60, LT-18170 Pabrade, Lithuania, and Intersurgical Ltd., Circuit House, Pitronnerie Road, Industrial Estate, St Peter Port, Guermsey, GY1 do hereby declare that

ECHIPAMED PLUS SRL str. Valea Trandafirilor 24 "B", of. 2-7 MD-2001, Chisinau Republic of Moldova

is our official distributor and local representative for the complete range of Intersurgical products, , in the territory of the Republic of Moldova.

We declare that above mentioned company is authorized to quote, sell, subsequently negotiate and sign contracts, perform installation and after sales service, as well as to perform all the procedures required for the expertise process at state registration (re-registration, etc.) of All Intersurgical Product Range of Medical Devices for Respiratory Support in the Republic Moldova, manufactured by us.

We hereby extend our full-warranty with respect to the Goods offered by the above company.

This authorization letter will remain valid until 31.12.2027 and all terms and conditions are as per the Distributor Agreement signed by both parties.

Yours sincerely

Stewen Williams

Global Sales Director



SURGICAL LTD.