

# EC Certificate - Production Quality Assurance

Directive 93/42/EEC on Medical Devices, Annex V

**No.** CE 698961  
**Issued To:** O & M Halyard, Inc.  
9120 Lockwood Blvd  
Mechanicsville  
Virginia  
23116  
USA

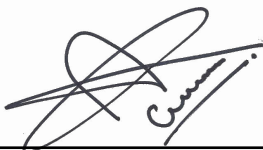
In respect of:

**The manufacture of Surgical Drapes.**

**Those aspects of Annex V related to securing and maintaining sterility in the manufacture of sterile surgical gowns, surgical drapes, surgical packs and examination gloves**

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex V. The quality assurance system meets the requirements of the directive. For the placing on the market of class IIb and class III products an Annex III certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 2797):



Albert Roossien, Regulatory Lead

First Issued: **2019-02-18**

Date: **2019-02-25**

Expiry Date: **2024-02-17**

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Page 1 of 2

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

This certificate was issued electronically and is bound by the conditions of the contract.

Information and Contact: BSI, Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands Tel: + 31 20 346 0780

BSI Group The Netherlands B.V. registered in The Netherlands under 33264284.

A member of BSI Group of Companies.

# EC Certificate - Production Quality Assurance

## Supplementary Information to CE 698961

Issued To:

**O & M Halyard, Inc.  
9120 Lockwood Blvd  
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Number	Device Name	Intended Purpose per IFU
<b>Class IIa</b>		
MD 0101	Transurethral Resection (T.U.R.) Drapes & Packs	N/A
<b>Class Is</b>		
MDS7006	Surgical Gowns	N/A
MDS7006	Surgical Drapes	N/A
MDS7006	Surgical Packs	N/A
MDS7006	Examination Gloves	N/A

First Issued: **2019-02-18**

Date: **2019-02-25**

Expiry Date: **2024-02-17**

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# EC Certificate - Production Quality Assurance

Directive 93/42/EEC on Medical Devices, Annex V

## List of Significant Subcontractors

Recognised as being involved in services relating to the product covered by:

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**9120 Lockwood Blvd**  
**Mechanicsville**  
**Virginia**  
**23116**  
**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Arc Royal Virginia Road Kells Co Meath Ireland	<b>EU Representative</b>
GRI Medical & Electronic Technology Co., Ltd 1805 Honggao Road Jiaxing Zhejiang 314031 China	<b>ETO Sterilization Manufacture</b>
Isomedix Operations, Inc. 1441 Don Haskins Drive El Paso Texas 79936 USA	<b>ETO Sterilization</b>

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**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
La Ada de Acuna S. De. R.L. De C.V. Av. Hidalgo No. 6 Esq., Blvd. Luis Donaldo Colosio Col. Educativa, Nogales Sonora 84093 Mexico	<b>Manufacture</b>
Lianyungang Aiyeh Non-Woven Products Co., Ltd No. 9 YunYang Rd. Huangjiuni Export Processing Zone Lianyungang, Jiangsu 222047 China	<b>Manufacture</b>
Master & Frank (Pinghu) Ent. Co., Ltd. No. 2000, Xingping II Rd. Pinghu Economic Development Zone Zhejiang P.R. China	<b>Manufacture</b>

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**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
O&M Halyard Honduras S.A. de C.V. Carretera Tegucigalpa Villanueva Cortes Honduras	<b>Manufacture</b>
O&M Halyard, Inc. 5405 Windward PKWY Alpharetta Georgia 3004 USA	<b>Regulatory Compliance</b>
SAFESKIN MEDICAL & SCIENTIFIC (THAILAND), LTD. 200 moo 8 Kanchanavanich Road Tambol Prik, Amphur Sadao Songkhla, 90120 Thailand	<b>Manufacture</b>

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**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Sterigenics S. de R. L. de C. V. James Watt No. 22 Parque Industrial Cuamatla Cuautitlan Izcalli Estado de México C.P. 54730 Mexico	<b>ETO Sterilization</b>
Sterigenics US, LLC 10821 Withers Cove Park Drive Charlotte North Carolina 28278 USA	<b>Gamma Irradiation</b>
Sterigenics US, LLC 1302 Avenue T Grand Prairie Texas 75050 USA	<b>ETO Sterilization</b>

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Directive 93/42/EEC on Medical Devices, Annex V

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**Virginia**  
**23116**  
**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Sterigenics US, LLC 687 S. Wanamaker Avenue Ontario California 91761 USA	<b>ETO Sterilization</b>
Sterigenics US, LLC 2971 Olympic Industrial Drive SE Suite 116 Atlanta Georgia 30339 USA	<b>ETO Sterilization</b>
Sterigenics US, LLC 2400 Airport Road Santa Teresa New Mexico 88008 USA	<b>ETO Sterilization</b>

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**23116**  
**USA**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Synergy Health (Thailand) Ltd 700/465 Amata Nakorn Industrial Estate Moo 7, Tambol Donhuaroh Amphur Muang Chonburi 20000 Thailand	<b>Gamma Sterilization</b>
Synergy Sterilisation (M) Sdn Bhd Plot 203 Kuala Ketil Industrial Estate Kuala Ketil Kedah 09300 Malaysia	<b>Gamma Sterilization</b>

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# EC Certificate - Production Quality Assurance Certificate History

Certificate No: **CE 698961**  
 Date: **2019-02-25**  
 Issued To: **O & M Halyard, Inc.**  
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**23116**  
**USA**

Date	Reference Number	Action
18 February 2019	9643055	First Issue.
Current	9643448	Traceable to NB 0086.

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Page 1 of 1

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.  
 This certificate was issued electronically and is bound by the conditions of the contract.

# Certificate of Registration

## QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

O & M Halyard, Inc.  
9120 Lockwood Blvd  
Mechanicsville  
Virginia  
23116  
USA

Holds Certificate No:

**FM 697013**

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

The design and development, manufacture and distribution of surgical gowns, protective garments, face masks, surgical drapes, orthopedic soft goods, patient care products, cold therapy products, C-Section packs, OB Packs, orthopedic packs, sterile and non-sterile examination gloves, Temperature management systems for the areas of general surgery and general medical use and sterilization wrap and non-woven materials for medical devices.

For and on behalf of BSI:

\_\_\_\_\_  
Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2014-12-09

Latest Revision Date: 2020-01-08

Effective Date: 2020-01-09

Expiry Date: 2023-01-08



Page: 1 of 3

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Certificate No: **FM 697013**

Location	Registered Activities
O & M Halyard, Inc. 9120 Lockwood Blvd Mechanicsville Virginia 23116 USA	Headquarter management activities.
O & M Halyard, Inc. 5405 Windward Parkway Alpharetta Georgia 30004 USA	The design and development of surgical gowns, protective garments, face masks, surgical drapes, orthopedic soft goods, patient care products, cold therapy products, C-Section packs, OB Packs, orthopedic packs, sterile and non-sterile examination gloves, Temperature management systems for the areas of general surgery and general medical use and sterilization wrap and non-woven materials for medical devices.
Halyard North Carolina, LLC 389 Clyde Fitzgerald Rd. Linwood North Carolina 27299 USA	The manufacture of nonwoven materials for medical devices, Sterilization wrap, and infection control products including disposable gowns and linens.
La Ada de Acuna 14 Finegan Road Del Rio Texas 78840 USA	Receiving and Incoming Inspection, Warehouse and Distribution.
O&M Halyard Honduras S.A. de C.V. Carretera Tegucigalpa Villanueva Cortes Honduras	The manufacture and distribution of disposable sterile and non-sterile surgical gowns.
La Ada de Acuna Avenida Hidalgo #16 Parque Industrial San Carlos Nogales Sonora 84092 Mexico	Receiving and incoming inspection. Manufacturer/Conversion of nonwoven materials.

Original Registration Date: 2014-12-09

Effective Date: 2020-01-09

Latest Revision Date: 2020-01-08

Expiry Date: 2023-01-08

Page: 2 of 3

This certificate remains the property of BSI and shall be returned immediately upon request.  
An electronic certificate can be authenticated [online](https://www.bsigroup.com/ClientDirectory). Printed copies can be validated at [www.bsigroup.com/ClientDirectory](https://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.

Certificate No: **FM 697013**

Location	Registered Activities
La Ada de Acuna Kim. 4.5 Carreterra Presa La Amistad Ciudad De Acuna Coahuila 26220 Mexico	The manufacture of non-sterile face masks (surgical isolation, industrial and respirator), non-surgical gowns, cold therapy products, and sterilization wrap.
La Ada de Acuna S.De. R.L. De C.V AV. Hidalgo #6 Esq., Blvd., Luis Donaldo Colosio, Col. Educativa Nogales Sonora 84093 Mexico	The manufacture of disposable products including sterile and non sterile surgical packs, gowns and components. The manufacture of temperature management systems for areas of general surgery.
Safeskin Medical & Scientific (Thailand) Ltd. 200 Moo 8, Kanchanavanich Road, Tambol Prik, Amphur Sadao, Songkhla 90120 Thailand	The design and development, production and distribution of industrial gloves, sterile and non-sterile examination gloves.



Original Registration Date: 2014-12-09

Effective Date: 2020-01-09

Latest Revision Date: 2020-01-08

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Page: 3 of 3

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# EC Certificate - Production Quality Assurance

Directive 93/42/EEC on Medical Devices, Annex V

**No.** CE 540596  
**Issued To:** **Teleflex Medical**  
**IDA Business and Technology Park**  
**Dublin Road**  
**Athlone**  
**Co. Westmeath**  
**Ireland**

In respect of:

**Those aspects of Annex V relating to securing and maintaining sterility in the manufacture of non-active respiratory, non-active gynaecological, non-active regional anaesthesia, non-active surgical and non-active urology devices.**

**Those aspects of manufacturing relating to obtaining sterility in the assembly of procedure packs in accordance with Article 12 of the Medical Devices Directive.**

**The manufacture of non-active and active surgical devices for adult and paediatric intraosseous infusion, bone marrow aspiration, bone marrow biopsy, bone lesion biopsy and non-active sterile urology catheters.**

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex V. The quality assurance system meets the requirements of the directive. For the placing on the market of class IIb and class III products an Annex III certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 2797):



Gary E Slack, Senior Vice President Medical Devices

First Issued: **2009-01-13**

Date: **2020-06-09**

Expiry Date: **2024-05-26**

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Page 1 of 3

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This certificate was issued electronically and is bound by the conditions of the contract.

# EC Certificate - Production Quality Assurance

## Supplementary Information to CE 540596

Issued To:

**Teleflex Medical  
IDA Business and Technology Park  
Dublin Road  
Athlone  
Co. Westmeath  
Ireland**

Number	Device Name	Intended purpose per IFU
<b>Class IIa</b>		
MD 0102	Sterile Intraosseous Vascular Access System	--
MD 1104	Non-sterile Intraosseous Vascular Access System	
MD 0102	Sterile Powered Bone Access	--
MD 1104	Non-sterile Powered Bone Access	
MD 0102	Sterile Sternal Intraosseous Device	--
MD 0101	Sterile Silicone Foley Catheter	--

First Issued: **2009-01-13**

Date: **2020-06-09**

Expiry Date: **2024-05-26**

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Page 2 of 3

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Information and Contact: BSI, Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands Tel: + 31 20 346 0780

BSI Group The Netherlands B.V. registered in The Netherlands under 33264284.

A member of BSI Group of Companies.



# EC Certificate - Production Quality Assurance

## Supplementary Information to CE 540596

Issued To:

**Teleflex Medical**  
**IDA Business and Technology Park**  
**Dublin Road**  
**Athlone**  
**Co. Westmeath**  
**Ireland**

Number	Device Name	Intended purpose per IFU
<b>Class Is</b>		
MD 0301	Intraosseous Vascular Access System Stabilizer	--
MD 0102	Powered bone access connector	--
MD 0101	Tracheostomy Tube Accessories	--
MD 0102	Tuohy Borst Adaptor	--
MD 0102	Syringe	--
MD 0101	Urology Dilator	--
MD 0101	Guedel Airway	--
MD 0101	Intrauterine Catheter Set	--
MD 0101	Sterile Container	--
MD 0101	Neckband	--
<b>Sterility aspects only</b>		
---	Procedure Packs under article 12	---

First Issued: **2009-01-13**

Date: **2020-06-09**

Expiry Date: **2024-05-26**

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# EC Certificate - Production Quality Assurance

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**Ireland**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
ArcRoyal Virginia Road Kells, Co. Meath Ireland	<b>Manufacture</b>
Arriol International Corporation Carretera San Isidro KM 17 Zona Franca San Isidro Santo Domingo Este Dominican Republic	<b>ETO Sterilization Manufacture</b>
Arrow International CR, a.s. Jamska 2359/47 Zdar Nad Sazavou 59101 Czech Republic	<b>Manufacture</b>
BBF Sterilisationservice GmbH Willy-Rüsch-Straße 10/1 71394 Kernern Germany	<b>Radiation (Gamma Sterilization)</b>

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**Co. Westmeath**  
**Ireland**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
CeMed GmbH Im Oberdorf 41 72419 Neufra Germany	<b>Assembly Packaging</b>
China Biotech Corporation No. 10, 33 rd., Road, Taichung Industrial Park Taichung Taiwan	<b>Radiation (Gamma Sterilization)</b>
Degania Silicone Limited Kibbutz 1513000 Degania Bet Israel	<b>Manufacture</b>
Donatelle Plastics, Inc. 501 County Road E-2 Extension New Brighton MN 55112 USA	<b>Manufacture</b>

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<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Foremount Enterprise Co., Ltd. No. 17, Alley 15, Lane 5 Shenan Street Shengang Dist 42944 Taichung City Taiwan	<b>Manufacture</b>
Iotron Industries USA 4394 East Park 30 Drive Columbia City Indiana 46725 USA	<b>Radiation (E Beam Sterilization)</b>
Medical Service GmbH Luisenstraße 8 75378 Bad Liebenzell/Unterhaugstett Germany	<b>Assembly Packaging</b>

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<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Mediplast Israel Ltd. 7 Hayarkon St. P.O. Box 13214 Industrial Zone Yavne 8122710 Israel	<b>ETO Sterilization</b>
Rose GmbH für Medizintechnik Gottbillstraße 25-30 54294 Trier Germany	<b>ETO Sterilization</b>
sfm medical devices GmbH Brückenstraße 5 63607 Wächtersbach Germany	<b>ETO Sterilization</b> <b>Manufacture</b>

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**Ireland**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Sparton Onyx, LLC 2920 Kelly Avenue Watertown South Dakota 57201-7249 USA	<b>Manufacture</b>
Sterigenics Germany GmbH Kasteler Straße 45 Wiesbaden 65203 Germany	<b>ETO Sterilization</b>
Sterigenics US, LLC 2400 Airport Road Santa Teresa New Mexico 88008 USA	<b>ETO Sterilization</b>

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**Ireland**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Steritec, Inc. P.O. Box 1969 1705 Enterprise Street Athens, TX 75751 United States of America	<b>ETO Sterilization</b>
Synergy Health Sterilisation UK Ltd 1 Alpha Court Capitol Park Thorne Doncaster DN8 5TZ United Kingdom	<b>ETO Sterilization</b>
Synergy Sterilisation (M) Sdn Bhd. Plot 203 Kuala Ketil Industrial Estate Kuala Ketil Kedah 09300 Malaysia	<b>ETO Sterilization</b>

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**Co. Westmeath**  
**Ireland**

<b>Subcontractor:</b>	<b>Service(s) supplied</b>
Teleflex Medical Sdn. Bhd. Lot PT 2577, Jalan Perusahaan 4 34600 Kamunting Perak Malaysia	<b>ETO Sterilization Manufacture</b>
Viant San Antonio, Inc. 7027 Fairgrounds Parkway San Antonio TX 78238 United States of America	<b>Manufacture</b>
Viant Upland, Inc. a.t.a. (formerly) Lake Region Medical 2052 West 11th Street Upland CA 91786 USA	<b>Manufacture</b>
Willy Rüsç GmbH Willy-Rüsç-Straße 4-10 71394 Kernen i.R., Germany	<b>Manufacture</b>

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# EC Certificate - Production Quality Assurance Certificate History

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 IDA Business and Technology Park  
 Dublin Road  
 Athlone  
 Co. Westmeath  
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Date	Reference Number	Action
13 January 2009	7245725	First issue.
17 March 2009	7325720	Company address amended. Extension to scope. Addition of Willy Rüsç, Germany as subcontractor for design and manufacture.
25 August 2009	7399908  7439096	Addition of SFM as significant subcontractor for manufacture. Addition of 'design' services supplied by Teleflex Medical, Malaysia, Arrow International CR, a.s. and Arrow International, Inc., Czech Republic. Correction of History page header. Intrauterine catheter added to scope.
08 September 2010	7558507	Scope reworded in accordance with generic device groups. Activity of 'Design' removed from all subcontractors and 'Control of Sterilisation' added. Certificate renewal.

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Date	Reference Number	Action
23 February 2011	7635647	Scope extended to include, 'Those aspects of manufacturing relating to securing and maintaining sterility in the assembly of procedure packs in accordance with Article 12 of the Medical Devices Directive.'  Addition of subcontractor, 'ArcRoyal Ltd., Virginia Road, Kells, Co. Meath, Ireland' for Manufacture and Control of Sterilization activities.
23 May 2012	7778468	Correction of significant subcontractor address.
04 February 2013	7932595	The addition of significant subcontractors Foremount Enterprise Co Ltd and Bidoia SAS Di Gianfranco Didia EC.
13 July 2015	8334933	Extension to scope to include 'The manufacture of non-active and active surgical devices for adult and paediatric intraosseous infusion, bone marrow aspiration, bone marrow biopsy and bone lesion biopsy.'  Significant subcontractor changes: Addition of Vidacare LLC, Lake Region Medical, Arriol International Corporation, Coastal Life Technologies, Inc & Sparton Onyx. LLC.

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 Dublin Road  
 Athlone  
 Co. Westmeath  
 Ireland**

Date	Reference Number	Action
28 August 2015	8406492	Certificate renewal. Removal from scope of 'those aspects of Annex V relating to securing and maintaining sterility in the manufacture of non-active digestive tract devices' and 'Those aspects of Annex V related to metrology in the manufacture of non-active respiratory devices'.
10 February 2016	8455693	Removal of Vidacare LLC from list of significant subcontractors. Service(s) supplied for Arriol International Corporation, Coastal Life Technologies Inc. and Lake Region Medical changed from crucial suppliers to Control of Sterilization, Manufacture. Service(s) supplied for Sparton Onyx. LLC changed from crucial supplier to Manufacture. Removal of repeated use of word 'devices' from scope.
28 July 2017	8762518	Change of address for Coastal Life Technologies. Addition of Donatelle Plastics Inc., 55112 New Brighton to list of significant subcontractors.
04 March 2019	7779566	Traceable to NB 0086.



# EC Certificate - Production Quality Assurance Certificate History

Certificate No: **CE 540596**  
 Date: **2020-06-09**  
 Issued To: **Teleflex Medical  
 IDA Business and Technology Park  
 Dublin Road  
 Athlone  
 Co. Westmeath  
 Ireland**

Date	Reference Number	Action
Current	3124053	<p>Certificate renewal.</p> <p>Addition of supplementary product information table.</p> <p>Update to scope to include non-active sterile urology catheters.</p> <p>Name change from Coastal Life Technologies to Viant San Antonio, Inc., Name change from Lake Region Medical to Viant Upland, Inc</p> <p>Removal of Control of Sterilization from Service(s) supplied for ArcRoyal Ltd., Arrow International CR, a.s. (Zdar), Viant San Antonio, Inc., Donatelle Plastics, Inc., Foremount Enterprise Co., Ltd., Viant Upland, Inc., sfm medical devices GmbH, Teleflex Medical Sdn. Bhd., and Willy Rüsç GmbH.</p> <p>Addition of ETO Sterilization to Service(s) supplied for sfm medical devices GmbH and Teleflex Medical Sdn. Bhd.</p> <p>Administrative correction of details for ArcRoyal, Arriol International Corporation, Arrow International CR, a.s., Donatelle Plastics, Inc., Foremount Enterprise Co., Ltd., Sparton Onyx. LLC, sfm medical devices GmbH, Teleflex Medical Sdn. Bhd. and Willy Rüsç GmbH.</p> <p>Removal of Arrow International CR a.s. (Hradec Kralove) and Bidoia SAS Di Gianfranco Didoia E.C.</p> <p>Addition of CeMed GmbH and Medical Service GmbH for Assembly and Packaging.</p>



# EC Certificate - Production Quality Assurance Certificate History

Certificate No: **CE 540596**  
 Date: **2020-06-09**  
 Issued To: **Teleflex Medical**  
**IDA Business and Technology Park**  
**Dublin Road**  
**Athlone**  
**Co. Westmeath**  
**Ireland**

Date	Reference Number	Action
	3124053	Addition of Degania Silicone Limited for Manufacture Addition of Steritec, Inc., Sterigenics US, LLC, Rose GmbH für Medizintechnik, Synergy Health Sterilisation UK Ltd, Sterigenics Germany GmbH, Mediplast Israel Ltd., and Synergy Sterilisation (M) Sdn Bhd. for ETO Sterilization Addition of Iotron Industries USA for E-beam Sterilization Addition of China Biotech Corporation and BBF Sterilisationservice GmbH for Gamma Sterilization.

# Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

Teleflex Medical  
IDA Business and Technology Park  
Dublin Road  
Athlone  
Westmeath  
Ireland

Holds Certificate No:

**FM 544574**

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

The design and manufacture of non-active digestive tract devices; non-active gynaecological devices, non-active regional anaesthesia devices, non-active respiratory devices, non-active surgical devices, non-active urology devices and active surgical devices.



For and on behalf of BSI:

\_\_\_\_\_  
Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2009-03-09

Latest Revision Date: 2020-02-12

Effective Date: 2020-02-12

Expiry Date: 2023-02-11

Page: 1 of 1



...making excellence a habit.™



The management system of

## Teleflex Medical

2917 Weck Drive, Research Triangle Park, NC, 27709, United States  
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 11 September 2018 until 14 July 2023  
and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 27 May 2021

Issue 29. Certified since 26 September 2000

Certification is based on reports numbered WWW/MC/06866

Multiple certificates have been issued for this scope  
The main certificate is numbered US97/10879.00

Authorised by

SGS United Kingdom Ltd, Notified Body 0120

202B Warrle Parkway, Weston-super-Mare, BS22 6WA UK  
t +44 (0)1934 522917 f +44 (0)1934 522137 www.sgs.com

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Page 1 of 2



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## Teleflex Medical

### Directive 93/42/EEC

on medical devices, Annex II (excluding section 4)

Issue 29

Detailed scope

Sterile Hem-o-lok Ligation Clips.  
Sterile Deknatel® PTFE pledgets.  
Sterile Polyester Nonabsorbable Surgical Sutures (POLYLENE/ "cottony"™ II, "silky" II POLYDEK®, TEVDEK® II, NextSitch®, Capio™, Fixt®, NiceLoop™, TEVDEK®).  
Sterile DEKLENE® II; DEKLENE® MAXXTM, CAPIOTM and FIXTM polypropylene non-absorbable surgical sutures.  
Sterile BONDEK® and BONDEK® Plus Polyglycolic Acid Synthetic Absorbable Surgical Sutures.  
Sterile Polyglytone 6211™ Monofilament Absorbable Surgical Sutures.  
Sterile MONODEK® Polydioxanone Absorbable Surgical Sutures.  
Sterile Hem-o-lok Automatic Clip Appliers.  
Metal Ligation System.

Sterile External stapling system (including stainless steel staples, staplers and removers), Sterile, Efx endo fascial closuresystem (abdominal access), Sterile, Efx shield fascial closure system (abdominal access), Sterile, Efx classic fascial closuresystem (abdominal access)  
Sterile stainless steel surgical Sutures  
Sterile FORCE FIBER® surgical sutures.  
Sterile Chest drainage and autotransfusion systems,  
Sterile Thoracic Catheters,  
Sterile and Non-sterile Aortic Punch,  
Non-sterile Self Retaining Tissue retractor/blades

Non-sterile Anaesthesia and respiratory Circuits including breathing bags and water traps,  
Non-sterile Heated Humidifiers, Non-sterile Non-Prefilled Humidifiers and Nebulizers, Non-sterile Small Volume Nebulizers, Sterile Prefilled Humidifiers and Nebulizers (saline or water) with adaptors, Sterile Prefilled unit dose vial /solution for nebulisation, Non-sterile Respiratory therapy Adaptors and connectors, Sterile Column and Reservoirs including adaptors, Non-sterile Nasal cannula (including gas sampling), Non-sterile Cannula and Supply Tubing, Nonsterile CPAP Cannula System, Non-sterile Manual resuscitators and PEEP valves, Non-sterile Respiratory and anaesthesia masks, Non-sterile Gas scavenging mask, Sterile Endotracheal tubes, Sterile Endobronchial tubes, Non-sterile Suction and Aspirating Tubes, Sterile Vented Thoracic Chest Seal, Sterile Operative Cholangiogram Catheters, Sterile Abdominal Access and Insufflation devices, Sterile Capillary drains, Sterile Percutaneous Surgical System (MiniLap and Grip graspers), Sterile Percutaneous Surgical System (Mini Polar electrosurgical probe and MiniGrip Bipolar Graspers), Percutaneous surgical System (Interchangeable electrosurgical tool tips) for laparoscopic surgery, Non-sterile Heat and Moisture Exchangers

Where the above scope includes class III medical device(s), a valid EC Design Examination Certificate according to Annex II (Section 4) is a mandatory requirement for each device in addition to this certificate to place that device on the market





**REF** CS-15802-E

**8** Fr. **2** Lumen **20** cm catheter length **.032** inch dia. spring-wire guide

## Two-Lumen Central Venous Catheterization Set with Blue FlexTip® Catheter

**Contents:**

- 1: Two-Lumen Indwelling Catheter: 8 Fr. x 20 cm Radiopaque Polyurethane with Blue FlexTip®, Extension Line Clamps
- 1: Spring-Wire Guide, Marked: .032" (0.81 mm) dia. x 23-5/8" (60 cm) (Straight Soft Tip on One End - "J" Tip on Other) with Arrow Advancer with ECG Mark
- 1: Catheter: 18 Ga. x 2-1/2" (6.35 cm) Radiopaque over 20 Ga. RW Introducer Needle
- 1: Introducer Needle: 18 Ga. x 2-1/2" (6.35 cm) XTW
- 1: Pressure Transduction Probe
- 1: Arrow® Raulerson Spring-Wire Introduction Syringe: 5 mL
- 1: Tissue Dilator: 9 Fr. (3.0 mm) x 10.2 cm
- 2: Dust Cap: Non-Vented
- 1: SecondSite™ Adjustable Hub: Fastener
- 1: SecondSite™ Adjustable Hub: Catheter Clamp

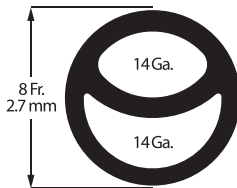
All components are CE 0086 unless otherwise noted.

**Rx only. Warning: Read all package insert warnings, precautions, and instructions prior to use. Failure to do so may result in severe patient injury or death.**

Not made with natural rubber latex.

Fluid path components are non-pyrogenic.

- fr** Jeu de cathétérisme veineux central à deux lumières avec cathéter Blue FlexTip
- de** Besteck für zweilumigen, zentralen Venenkatheter mit Blue-FlexTip-Katheter
- it** Set per cateterismo della vena centrale a due lumi con catetere Blue FlexTip
- pl** Dwukanałowy zestaw do cewnikowania żył centralnych z cewnikami Blue FlexTip
- pt** Conjunto de Cateterização Venosa Central com Lúmen Duplo e Cateter Blue FlexTip
- ru** Комплект для катетеризации главных вен с двухпросветным катетером Blue FlexTip
- sl** Komplet za centralno vensko katetrizacijo z dvema svetlinama s katetrom Blue FlexTip
- es** Conjunto de cateterización venosa central de dos luces con catéter Blue FlexTip
- sv** Set för tvåkanalig central venkatetrering med Blue FlexTip-kateter
- tr** İki Lümenli Santral Venöz Kateterizasyon Seti, Blue FlexTip Kateteri ile



Lumen	Priming Volume* (mL)	Flow Rate† (mL/hr)
Distal (14 Ga.)	0.75	5600
Proximal (14 Ga.)	0.80	5100

\* Priming volumes are approximate and are done without accessories.

† Flow rates are done with normal saline, room temperature, 100 cm head height and represent approximate flow capabilities.



LOT



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Co. Westmeath, Ireland



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Reading, PA 19605 USA

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LBL028056 (02/20)



**REF** CS-15802-E

**8** Fr. **2** Lumen **20** cm catheter length **.032** inch dia. spring-wire guide

**Two-Lumen Central Venous Catheterization Set with Blue FlexTip® Catheter**







REF CS-15853-E

8.5 Fr. 3 Lumen

20 cm catheter length .032 inch dia. spring-wire guide

# Three-Lumen CVC

**Contents:**

- 1: Three-Lumen Catheter: 8.5 Fr. (3.0 mm OD) x 20 cm
- 1: Spring-Wire Guide, Marked: .032" (0.81 mm) dia. x 23-5/8" (60 cm) (Straight Soft Tip on One End - "J" Tip on Other) with Arrow Advancer
- 1: Catheter: 18 Ga. x 2-1/2" (6.35 cm) Radiopaque over 20 Ga. RW Introducer Needle
- 1: Introducer Needle: 18 Ga. x 2-1/2" (6.35 cm) XTW
- 1: Pressure Transduction Probe
- 1: Arrow® Raulerson Spring-Wire Introduction Syringe: 5 mL
- 1: Tissue Dilator: 9 Fr. (3.0 mm) x 10.2 cm
- 3: Dust Cap: Non-Vented
- 1: SecondSite™ Adjustable Hub: Fastener
- 1: SecondSite™ Adjustable Hub: Catheter Clamp

All components are CE 2797 unless otherwise noted.

**Rx only.**

**Warning: Read all package insert warnings, precautions, and instructions prior to use. Failure to do so may result in severe patient injury or death. [www.teleflex.com/IFU](http://www.teleflex.com/IFU)**

**California Prop. 65**  
 ⚠ **WARNING:** Cancer and Reproductive Harm.  
[www.P65Warnings.ca.gov](http://www.P65Warnings.ca.gov)

Not made with natural rubber latex.  
 Fluid path components are non-pyrogenic.



Lumen	Priming Volume* (mL)	Gravity Flow Rate† (mL/hr)
Distal (16 Ga.)	0.47	2930
Medial (14 Ga.)	0.89	6267
Proximal (16 Ga.)	0.47	3731

\* Priming volumes are approximate and are done without accessories.

† Flow rate values are approximate and are determined using deionized water at 100 cm head height.

EU Authorized Representative and Importer:

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 Dublin Road, Athlone  
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 2797

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LBL055441 (2021-01)

LOT



8.5 Fr. 3 Lumen

20 cm catheter length .032 inch dia. spring-wire guide

REF CS-15853-E

# Three-Lumen CVC





REF CS-12703

7 Fr. 3 Lumen

16 cm catheter length

.032 inch dia. spring-wire guide

# Three-Lumen CVC

**Contents:**

- 1: Three-Lumen Catheter: 7 Fr. (2.5 mm OD) x 16 cm
- 1: Spring-Wire Guide, Marked: .032" (0.81 mm) dia. x 17-3/4" (45 cm) (Straight Soft Tip on One End - "J" Tip on Other) with Arrow Advancer
- 1: Catheter: 18 Ga. x 2-1/2" (6.35 cm) Radiopaque over 20 Ga. RW Introducer Needle
- 1: Introducer Needle: 18 Ga. x 2-1/2" (6.35 cm) XTW
- 1: Syringe: 5 mL Luer-Slip
- 1: Tissue Dilator: 8.5 Fr. (2.8 mm) x 10.2 cm
- 3: Dust Cap: Non-Vented
- 1: SecondSite™ Adjustable Hub: Fastener
- 1: SecondSite™ Adjustable Hub: Catheter Clamp

All components are CE 2797 unless otherwise noted.

**Rx only**

**Warning: Read all package insert warnings, precautions, and instructions prior to use. Failure to do so may result in severe patient injury or death. [www.teleflex.com/IFU](http://www.teleflex.com/IFU)**

**California Prop. 65**

**⚠ WARNING:** Cancer and Reproductive Harm. [www.P65Warnings.ca.gov](http://www.P65Warnings.ca.gov)

**Not made with natural rubber latex.**

Fluid path components are non-pyrogenic.



Lumen	Priming Volume* (mL)	Gravity Flow Rate† (mL/hr)
Distal (16 Ga.)	0.4	2947
Medial (18 Ga.)	0.4	1552
Proximal (18 Ga.)	0.4	1702

\* Priming volumes are approximate and are done without accessories.

† Flow rate values are approximate and are determined using deionized water at 100 cm head height.

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LBL055929 (2021-03)

LOT



7 Fr. 3 Lumen

16 cm catheter length

.032 inch dia. spring-wire guide

REF CS-12703

# Three-Lumen CVC



REF CS-15854-E

8.5 Fr. 4 Lumen

20 cm catheter length

.032 inch dia. spring-wire guide

# Four-Lumen CVC

**Contents:**

- 1: Four-Lumen Catheter: 8.5 Fr. (3.0 mm OD) x 20 cm
- 1: Spring-Wire Guide, Marked: .032" (0.81 mm) dia. x 23-5/8" (60 cm) (Straight Soft Tip on One End - "J" Tip on Other) with Arrow Advancer with ECG Mark
- 1: Catheter: 18 Ga. x 2-1/2" (6.35 cm) Radiopaque over 20 Ga. RW Introducer Needle
- 1: Introducer Needle: 18 Ga. x 2-1/2" (6.35 cm) XTW
- 1: Pressure Transduction Probe
- 1: Syringe: 5 mL Luer-Slip
- 1: Arrow® Raulerson Spring-Wire Introduction Syringe: 5 mL
- 1: Tissue Dilator: 9 Fr. (3.0 mm) x 10.2 cm
- 4: Dust Cap: Non-Vented
- 1: SecondSite™ Adjustable Hub: Fastener
- 1: SecondSite™ Adjustable Hub: Catheter Clamp

**Warning: Read all package insert warnings, precautions, and instructions prior to use. Failure to do so may result in severe patient injury or death. [www.teleflex.com/IFU](http://www.teleflex.com/IFU)**  
**Not made with natural rubber latex.**  
 Fluid path components are non-pyrogenic.

All components are CE 2797 unless otherwise noted.



Lumen	Priming Volume* (mL)	Gravity Flow Rate† (mL/hr)
Distal (16 Ga.)	0.5	2399
Medial 1 (14 Ga.)	0.6	4200
Medial 2 (18 Ga.)	0.4	1104
Proximal (18 Ga.)	0.4	1204

\* Priming volumes are approximate and are done without accessories.

† Flow rate values are approximate and are determined using deionized water at 100 cm head height.

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LBL055990 (2021-04)

LOT



8.5 Fr. 4 Lumen

20 cm catheter length

.032 inch dia. spring-wire guide

REF CS-15854-E

# Four-Lumen CVC



REF CS-16553-E

5.5 Fr. 3 Lumen 13 cm catheter length .018 inch dia. spring-wire guide

# Pediatric Three-Lumen CVC

**Contents:**

- 1: Three-Lumen Catheter: 5.5 Fr. (1.9 mm OD) x 13 cm
- 1: Spring-Wire Guide: .018" (0.46 mm) dia. x 17-3/4" (45 cm) (Straight Soft Tip on One End - "J" Tip on Other)
- 1: Introducer Needle: 21 Ga. x 1-1/2" (3.81 cm) TW
- 1: Catheter: 22 Ga. x 1-3/4" (4.45 cm) Radiopaque over 25 Ga. RW Introducer Needle with 5 mL Luer-Slip Syringe
- 1: Tissue Dilator: 6 Fr. (2.0 mm) x 7.6 cm
- 1: Tissue Dilator: 6 Fr. (2.0 mm) x 4.1 cm
- 3: Dust Cap: Non-Vented
- 1: SecondSite™ Adjustable Hub: Fastener
- 1: SecondSite™ Adjustable Hub: Catheter Clamp

All components are CE 2797 unless otherwise noted.

**Rx only**

**Warning: Read all package insert warnings, precautions, and instructions prior to use. Failure to do so may result in severe patient injury or death. [www.teleflex.com/IFU](http://www.teleflex.com/IFU)**

**Not made with natural rubber latex.**

Fluid path components are non-pyrogenic.



Lumen	Priming Volume* (mL)	Gravity Flow Rate† (mL/hr)
Distal (20 Ga.)	0.32	988
Medial (22 Ga.)	0.38	370
Proximal (22 Ga.)	0.34	422

\* Priming volumes are approximate and are done without accessories.

† Flow rate values are approximate and are determined using deionized water at 100 cm head height.

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LBL056000 (2021-03)

LOT



REF CS-16553-E

5.5 Fr. 3 Lumen 13 cm catheter length .018 inch dia. spring-wire guide

# Pediatric Three-Lumen CVC