



Directive 93/42/EEC on Medical Devices, Annex II Section 4

No. CE 69002

Issued To: Cordis Corporation

14201 North West 60th Avenue

Miami Lakes

Florida 33014 USA

In respect of:

Cordis 6F 0.070" Vista Brite Tip® Guiding Catheters

BSI has performed a design examination on the above devices in accordance with the Council Directive 93/42/EEC, Annex II Section 4. The design conforms to the requirements of this directive. For marketing of these products an additional Annex II excluding Section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

Stewart Brain, Head of Compliance & Risk -

Medical Devices

First Issued: **2002-08-19** Date: **2017-07-31** Expiry Date: **2022-08-18**

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Page 1 of 4

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.





Supplementary Information to CE 69002

Issued To: Cordis Corporation

14201 North West 60th Avenue

Miami Lakes Florida 33014 USA

Product:

General Designation: 670-XXX-XX

XXX - XXX - XXX 123 456 789

Number / Designation	Limitation
1 Outer Diameter (last digit of French Size)	Will always be 6 French
2,3 Lumen Size (last 2 digits in thousandths of an inch)	0.065-0.075 inches
4,5,6 Configuration	
000-299 Standard (subassembly) design	Odd numbers contain a side hole
300-599 Design variation 1	Even numbers do not contain a side
600-899 Design variation 2	hole
900-999 Design variation 3 / Overflow	
7,8 Length (last 2 digits in cm)	50-125 cm
In addition digits 8 or 9 may contain a single letter	
code.	
For example,	(2)
E – Econopack	
L – Long Bright Tip	
N – Guiding catheter with an introducer	ESSE

Modified Standards: SMXXXX and SMXXXXX

First Issued: **2002-08-19** Date: **2017-07-31** Expiry Date: **2022-08-18**

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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.





Supplementary Information to CE 69002

Issued To:

Cordis Corporation 14201 North West 60th Avenue Miami Lakes Florida 33014 USA

Certificate History

Date	Reference Number	Action
19 August 2002	EQ 10040583	First issue.
		Change of format of the Certificate product listing.
09 December 2003	EQ 10052521	The addition of Roden, The Netherlands to the list of Sterilization companies used. New format for the presentation of the catalogue numbers.
23 August 2004	EQ 10059974	Shelf Life extension to 3 years and revision of history to remove items prior to first issue under CE 69002.
31 March 2006	EQ 10078108	Changes to wildcards.
08 June 2007	EQ 10088941	Change in pouch heat seal coating from CR 27 (ex. Perfecseal) and 703 HSC (ex. Mangar) to PTH 025 (ex. Mangar).
01 August 2007	EQ 10089895	Add pouch with RLE004 PET/PE film PTH 034 heat seal coating, and Tyvek 1073B.
31 August 2007	EQ 10089893	Certificate renewal.
16 August 2012	10136333	Certificate renewal
		Modified Standards: SMXXXX and SMXXXXX added as they were inadvertently omitted when CE 69002 was split from CE 01110 under EQ 10040583.

First Issued: **2002-08-19** Date: **2017-07-31** Expiry Date: **2022-08-18**

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Supplementary Information to CE 69002

Issued To:

Cordis Corporation

14201 North West 60th Avenue

Miami Lakes Florida 33014 USA

Certificate History

Date	Reference Number	Action
04 February 2016	10160465	Change affecting DuPont Tyvek 1073B packaging material – all product codes are affected.
Current	8763074	Certificate Renewal. Removed Envoy® Guiding Catheter from certificate scope and product catalogue.

First Issued: **2002-08-19** Date: **2017-07-31** Expiry Date: **2022-08-18**

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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.





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No.

Issued To: Teleflex Medical

IDA Business and Technology Park

Dublin Road Athlone

CE 540595

Co. Westmeath

Ireland

In respect of:

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

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

Frank Lee, EMEA Compliance & Risk Director

First Issued: 13 January 2009 Date: 28 August 2015 Expiry Date: 07 September 2020

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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.



72419 Neufra Germany



EC Certificate - Full Quality Assurance System

Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 540595**

Date: 28 August 2015
Issued To: Teleflex Medical

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Subcontractor:	Service(s) supplied				
Arrow International CR, a.s. Jamska 2359/47 59101 Zdar nad Sazavou Czech Republic	Control of Sterilization Design Manufacture				
Arrow International CR, a.s. Prazska 209 50004 Hradec Kralove Czech Republic	Control of Sterilization Design Manufacture				
Arrow Medical Ltd Hatton Gardens Industrial Estate Kington HR5 3RB United Kingdom	Crucial Supplier	QUA			
CeMed GmbH Oberdorf 41	Control of Sterilization Manufacture				





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 540595**

Date: 28 August 2015 Issued To: **Teleflex Medical**

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Subcontractor:

Service(s) supplied

Chelle Medical Limited

PO Box 221 Le Rocher Victoria Mahe

Seychelles

Crucial Supplier

Forefront (Xiamen) Medical

Devices Co., Ltd

No 26 & 28 Haijing Dong Lu Haicang Xiamen Export Processing Zone 361026, Xiamen, Fujian

China

Crucial Supplier

Forefront Medical Technology Pte Ltd 35 Joo Koon Circle, 6th Floor Singapore 629110 Singapore

Crucial Supplier





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 540595**

Date: 28 August 2015
Issued To: Teleflex Medical

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Subcontractor:

Service(s) supplied

Crucial Supplier

M.E.M., Inc. 8 Bishop Lane Madison

Connecticut 06443

USA

Crucial Supplier

Parker Medical Systems Division -Merrillville 1201 East 86th Place Merrillville Indiana 46410 USA

Plaxtron Industrial (M) Sdn. Bhd. Plot 28, Kawasan Perusahaan Jelapang II Zon Perdagangan Bebas 30020 Ipoh Perak Malaysia **Crucial Supplier**



049909

Singapore



EC Certificate - Full Quality Assurance System

Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 540595**

Date: 28 August 2015
Issued To: Teleflex Medical

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Subcontractor:	Service(s) supplied	
SP Medical A/S Møllevej 1 4653 Karise Denmark	Control of Sterilization Design Manufacture	
Süddeutsche Feinmechanik GmbH (SFM) Brückenstrasse 5 D-63607 Wächtersbach Germany	Control of Sterilization Manufacture	
Teleflex Medical Sdn. Bhd. Lot PT2577, Jalan Perusahaan 4 34600 Kamunting Perak Malaysia	Control of Sterilization Design Manufacture	PQUA
Teleflex Medical Asia Pte. Ltd. 6 Battery Road #07-02	Control of Sterilization Design	

Manufacture





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 540595**

Date: 28 August 2015 Issued To: **Teleflex Medical**

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Subcontractor:

Service(s) supplied

The Laryngeal Mask Company (Malaysia) Sdn. Bhd. Lot 19 & 1920 Industrial Zone Phase 1 Kulim Hi-Tech Park, Kulim 09000 Malaysia

Crucial Supplier

Tianiin Medis Medical Device Co. Ltd 10A Tianzhi Industrial Centre No 12 Hong Yuan Road Xiqing Economic Development Area

300385 Tianjin City

China

Control of Sterilization Manufacture

Willy Rüsch GmbH Willy Rüsch-Strasse 4-10 D-71394 Kernen Germany

Control of Sterilization Design **Manufacture**





Certificate No:

CE 540595

Date:

28 August 2015

Issued To:

Teleflex Medical

IDA Business and Technology Park

Dublin Road

Athlone

Co. Westmeath

Ireland

Date	Reference Number	Action				
13 January 2009	7245725	First issue				
17 March 2009	7325719	Company address amended. Extension to scope. Addition of Willy Rüsch, Germany as subcontractor for design and manufacture				
25 August 2009	7399879	Addition of 'epidural catheter Epistar and Epistar CSE' to scope. Addition of SFM as significant subcontractor for manufacture. Addition of 'design' to services supplied by Teleflex Medical Malaysia, Arrow International CR, a.s. and Arrow International Inc., Czech Republic				
11 November 2009	7455515	Addition of CeMed GmbH for manufacturing to the list of significant subcontractors				
20 April 2010	7497906	Laryngeal Mask added to scope. Addition of Tianjin Medis Medical Device Co. Ltd as significant subcontractor for manufacture				
08 September 2010	7558508	Scope reworded in accordance with generic device groups. Certificate renewal				
23 May 2012	7778467	Correction of significant subcontractor address and addition of new scope activities for subcontractors				

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Page 1 of 2

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Certificate No:

CE 540595

Date:

28 August 2015

Issued To:

Teleflex Medical

IDA Business and Technology Park

Dublin Road Athlone

Co. Westmeath

Ireland

Date	Reference Number	Action
04 February 2013	7932588	The addition of a significant subcontractor SP Medical A/S
14 May 2014	8134266	Addition of peripheral angioplasty balloon catheters to product family, covered by scope expression 'non-active surgical devices'.
		Addition of significant subcontractors Hotspur Technologies, Inc and Teleflex Medical Asia Pte Ltd
09 March 2015	8293488	Addition of 8 crucial suppliers
28 August 2015	8406490	Certificate renewal.
		Removal of Hotspur Technologies, Inc. from list of significant subcontractors.

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Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No. CE 510108

Issued To: Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

In respect of:

The design, development and manufacture of coronary and peripheral dilation catheters, stent systems, including covered stents, drug eluting stents, Bioresorbable Vascular Scaffold (BVS) Systems, carotid and peripheral stent systems, embolic protection systems, femoral vessel closure devices and the related instruments necessary for the deployment of the closure devices, guidewires, mitral valve repair systems, and associated accessories.

Those aspects of Annex II related to securing and maintaining the sterility of guide wire extensions, torque devices, hemostatic valves, introducers and flushing tools.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

Stewart Brain, Head of Compliance & Risk -

Medical Devices

First Issued: **2006-08-01** Date: **2017-12-22** Expiry Date: **2020-10-16**

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Page 1 of 1

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Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 510108**Date: **2017-12-22**

Ireland

Issued To: Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Subcontractor: Service(s) supplied

Abbott Ireland **ETO Sterilization**Ballytivnan
Sligo

Abbott Vascular International BVBA **EU Representative**

Park Lane Culliganlaan, 2B 1831 Diegem Belgium

Abbott Vascular Netherlands B.V.

Argonstraat 1

6422 PH Heerlen

The Netherlands

Distribution

Labelling

Packaging

Abbott Vascular
26531 Ynez Road
Temecula
California 92591
USA

Design
Development
E beam Sterilization
Manufacture





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 510108**

Date: **2017-12-22**

Issued To: Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Subcontractor:

Service(s) supplied

Abbott Vascular 3885 Bohannon Drive

Menlo Park CA 94025 USA Design
Development
Distribution
Manufacture

Abbott Vascular

52 Calle, 3, B31, Coyol Free Zone

El Coyol Alajuela Costa Rica Manufacture

Abbott Vascular

Building PR-17, Road #2 km. 58.0

Cruce Davila Barceloneta 00617 Puerto Rico **Manufacture**

Abbott Vascular Cashel Road Clonmel Tipperary

Ireland

Design Development Manufacture





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 510108**

Date: **2017-12-22**

Issued To: Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Subcontractor:

Service(s) supplied

Abbott West Distribution Center 42301 Zevo Drive

42301 Zevo Drive Temecula California 92590 USA Distribution Manufacture

Acme Monaco 75 Winchell Drive New Britain CT 06052

USA

Manufacture

Ad)medes Schuessler GmbH Rastatter Strasse 15 75179 Pforzheim Germany **Manufacture**





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No:

CE 510108

Date:

2017-12-22

Issued To:

Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 **USA**

Subcontractor:

Service(s) supplied

Availmed S.A. de C.V. C. Industrial Lt. 001 Mz.105 No. 20905 Int. A

Col. Cd. Industrial

Tijuana

Baja California

22444 Mexico **Manufacture**

Nitinol Devices and Components, Inc.

Costa Rica, S.R.L Coyol Free Zone Building B14 and B15 El Coyol, Alajuela

Manufacture

Nitinol Devices and Components, Inc

47533 Westinghouse Drive

Fremont CA 94539 **USA**

Costa Rica

Manufacture





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No:

CE 510108

Date:

2017-12-22

Issued To:

Abbott Vascular 3200 Lakeside Drive

Santa Clara

California 95054 USA

Subcontractor:

Service(s) supplied

Novartis Pharma AG Lichtstrasse 35

Lichtstrasse 35 Basel

CH-4056 Switzerland **Crucial Supplier**

Parter Sterilization Services LLC 17115 Kingsview Ave

Carson CA 90746 USA **ETO Sterilization**

Rose Technologies 1440 Front Avenue NW Grand Rapids Michigan 49504 USA **Manufacture**





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No:

CE 510108

Date:

2017-12-22

Issued To:

Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Subcontractor:

Service(s) supplied

Sterigenics Costa Rica S.R.L. Zona Franca PROPARK Calle Principal, Edificio 10

El Coyol Alajuela Costa Rica ETO Sterilization

Sterigenics Germany GmbH Kasteler Strasse 45

65203 Wiesbaden

Germany

ETO Sterilization

Sterigenics UK Limited Cotes Park Estate Somercotes Alfreton DE55 4NJ United Kingdom **ETO Sterilization**





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No: **CE 510108**

Issued To: Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

2017-12-22

Subcontractor:

Date:

Service(s) supplied

Sterigenics US, LLC 2400 Airport Road Santa Teresa New Mexico 88008 USA **ETO Sterilization**

Sterigenics US, LLC 4900 South Gifford Avenue Los Angeles CA 90058 USA **ETO Sterilization**

Sterigenics US, LLC 7695 Formula Place San Diego California 92121 USA E beam Sterilization





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

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

Certificate No:

CE 510108

Date:

2017-12-22

Issued To:

Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Subcontractor:

Service(s) supplied

Synergy Health AST, SRL B16, Street 4, Avenue 0 El Coyol Free Zone 20102 El Coyol Alajuela Costa Rica **E beam Sterilization**

Synergy Health Ireland Ltd. IDA Business & Technology Park Sragh Industrial Estate Tullamore, Co. Offaly Ireland E beam Sterilization ETO Sterilization

Teleflex Medical OEM 50 Plantation Drive Jaffrey NH 03452

USA

Manufacture





Certificate No: **CE 510108**

Date: **2017-12-22**

Issued To: Abbott Vascular 3200 Lakeside Drive

Santa Clara California

95054 USA

Date	Reference Number	Action
01 August 2006	4068482	First Issue based on CE 00946.
13 March 2007	4941821	Isotron Ireland, Ltd added to the list of significant subcontractors.
15 November 2007	7104034	Addition of Abbott Ireland (Galway) to the list of significant subcontractors. Addition of design and development of services supplied by Temecula.
01 August 2008	7200338	Addition of Abbott Vascular, Murrieta and Abbott Vascular, Barceloneta to list of significant subcontractors for manufacturing activities. Removal of Abbott Vascular, Dorado facility.
		Transfer of product families from Abbott Vascular, Vascular Solutions FQA certificate CE 525963.
18 February 2009	7292729	Remove Business Unit name (Cardiac Therapies) from the 'issued to' address and the Abbott Vascular, Murrieta facility address in the list of subcontractors.
		Addition of AD)MEDES Schuessler GmbH to list of significant subcontractors for manufacturing activities.
20 April 2010	7510769	Addition of Creganna-Tactx Medical to list of significant subcontractors for manufacturing activities and addition of Abbott Vascular International BVBA as EU Authorized Representative.

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Page 1 of 4

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Certificate No:

CE 510108

Date:

2017-12-22

Issued To:

Abbott Vascular

3200 Lakeside Drive

Santa Clara California 95054 USA

Date	Reference Number	Action				
		Renewal of certification				
		Removal of Sterigenics (Salt Lake City), Abbott Ireland (Galway) and Isotron Ireland as significant subcontractors. Remove Abbott Vascular Sterilization from Clonmel manufacturing site.				
12 October 2010	7581791	Addition of Sterigenics (New Mexico) as significant subcontractor.				
		Removal of atherctomy catheters and motor drive units from the scope. Redefine stents as stent systems.				
		Addition of Abbott West Distribution Center and Abbott Vascular Devices Holland B.V. as a significant subcontractor.				
10 November 2011	7765633	Addition of LEONI Studer Hard AG to list of significant subcontractors for E beam sterilization.				
13 December 2011	7766500	Addition of the Abbott Vascular Manufacturing Site in Alajuela, Costa Rica as a significant subcontractor.				
31 May 2012	7804693	Addition of Synergy Health Ireland Ltd as a significant subcontractor for e-beam sterilization. Name of subcontractor Abbott Vascular Devices Holland B.V. changed to Abbott Vascular Netherlands B.V. and address updated. Administrative changes on certificate.				
19 September 2012	7903213	Addition of Accellent as significant subcontractor for TREK family. Addition of Abbott Vascular Costa Rica Main Building as significant subcontractor for manufacturing.				

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Page 2 of 4

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Certificate No: **CE 510108**

Date: **2017-12-22**

Issued To: Abbott Vascular 3200 Lakeside Drive

Santa Clara California 95054

USA

Date	Reference Number	Action				
21 December 2012	7911227	Addition of Abbott (Nutritional) Ireland Sligo to the list of significant subcontractors for the sterilization.				
		Scope updated to include "including covered stents".				
02 July 2013	7991114	Removal of Abbott Vascular - Alajuela Costa Rica, as a significant subcontactor. Change name of subcontractor from LEONI Studer Hard AG to LEONI Studer AG. Reclassify Funnel Introducer, Guide Wire Introducer, Duostat Rotating Hemostatic Valve, Rotating Hemostatic Valve, Guide Wire Introducer Accessory Kit and Guide Wire Accessory Kit with CoPilot from Class II a to Class I (Sterile).				
May 28, 2014	8164752	Addition of NovoSci and Sterigenics in Wiesbaden for the service of ETO sterilization, Synergy Health in Costa Rica for the service of Ebeam sterilization and Availmed S.A. de C.V. for service of manufacturer due to several product transfers.				
05 February 2015	8268209	Update to add Drug Eluting Stents to the scope. Addition of significant subcontractors OK International, LTD and Sterigenics UK Limited.				
31 March 2015	8283470	Addition of Vessel Closure Devices to the scope of certification as part of a transfer from the Abbott Vascular Redwood City facility. Addition of significant subcontractors Teleflex Medical and Acme Monoco for manufacture and Synergy Health Ireland Ltd for EO Sterilization.				

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Page 3 of 4

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Certificate No: **CE 510108**

Date: **2017-12-22**

Issued To: Abbott Vascular 3200 Lakeside Drive

Santa Clara California 95054

9505 USA

Date	Reference Number	Action				
13 April 2015	8296689	Addition of Bioresorbable Vascular Scaffold (BVS) Systems to the scope of certification.				
08 July 2015	8359594	Addition of Sterigenics Costa Rica S.R.L. as a significant subcontractor for ETO sterilization.				
07 September 2015	8411826	Renewal of certification. Removal of subcontractors: Accellent, Inc., Creganna, NovoSci Corp and OK International, LTD. Removal of Abbott Vascular Murrieta site: facility closed down. Typo correction (LEONI Studer AG address, Sterigenics names).				
19 December 2015	8427566	Scope extension to include the MitraClip NT System under Abbott Vascular's Quality System.				
13 July 2016	8558860	Removal of "coronary and peripheral guiding catheters" from scope of certification and the addition of Availmed S.A. de C.V. Baja California location as significant subcontractor.				
Current	8863184	Scope change from "Arterial" to "Femoral" for vessel closure devices. Removal of Availmed in La Mesa, Tijuana, Mexico for manufacturing services, and LEONI in Switzerland for Ebeam Sterilization. Addition of NOVARTIS as a crucial supplier. Add design and development services to Abbott in Clonmel, Ireland.				

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TAM KALİTE GÜVENCE BELGESİ

MERIL LIFE SCIENCES PVT. LTD.

firması

MUKTANAND MARG, CHALA, VAPI-396191. GUJARAT, HİNDİSTAN

adresinde

SİROLİMUS SALINIMLI BİYOEMİLEBİLİR VASKÜLER SCAFFOLD SİSTEMİ (MERES100TM LINEAGE)

kapsamı için

93/42/AT – Tıbbi Cihaz Yönetmeliği Tam Kalite Güvence Sistemi EK-II (Bölüm 4 Hariç)

gereklerine göre tetkik edilmiş ve belgelendirilmiştir.

Onaylanmış Kuruluş Numarası:

1783

Belge Veriliş Tarihi:

17.05.2019

Geçerlilik Tarihi:

17.05.2024

GMDN Kodu:

56304

Sınıflandırma:

Smif III

AT Tasarım İnceleme Belgesi Numarası:

1783-MDD-119

İnceleme Rapor Numarası:

1542-MDD-099/2018-01

Belge Değişiklik Tarihi / Nedeni:

Tıbbi Cihaz Yönetmeliği Ek II Bölüm 3'e göre Kalite Sisteminin Teknik Düzenleme / Uyumlaştırılmış Standard gereklerini karşıladığını gösteren, işbu belge ile Kuruluş; tetkiki yapılan kalite sistemi kapsamında, CE Uygunluk İşaretini, aşağıda gösterildiği şekilde iliştirme ve Onaylanmış Kuruluş numarasını kullanmaya yetkilidir. Onaylanmış Kuruluş Tıbbi Cihaz Yönetmeliğinin EK II, 5. bölümüne istinaden gerekli gözetimleri yapına hakkına sahiptir.

Bu belge kapsamında bulunan Sınıf III ürün(ler)ün CE işaretlemesi için, Tıbbi Cihaz Yönetmeliği EK-II, 4. Bölümüne göre düzenlenen Tasarım İnceleme Belgesi de gerekmektedir.

CE

Belge No: 1783 -MDD- 118

TANDARTLARI ENGLISION OF THE PROPERTY OF THE P

SOF

SEZAİ DOĞAN Direktifler Müdürü ANKARA Rev 00, 17/05/2019

Bu belge ancak TSE-Onaylanmış Kuruluş Numarası 1783 mührü ile geçerlidir.



CERTIFICATE OF FULL QUALITY ASSURANCE

MERIL LIFE SCIENCES PVT LTD

located at the address
MUKTANAND MARG CHALA VAPI-396191 GUJARAT INDIA

for the scope of

SIROLIMUS ELUTING BIORESORBABLE VASCULAR SCAFFOLD SYSTEM (MERES100TM LINEAGE)

has been examined and certified to the requirements of

93/42/EEC – Medical Device Directive Full Quality Assurance System Annex II (Excluding Section 4)

Notified Body Number:

1783

Date of Issue:

17.05.2019

Valid Until:

17.05.2024

GMDN Code:

56304

Classification:

Class III

EC Design Examination Certificate Number:

1783-MDD-119

Inspection Report Number:

1542-MDD-099/2018-01

Date / Reason of the Certificate Revision:

This certificate remarks that quality system meets requirements of the technical regulations / harmonized standards according to Medical Device Directive Annex II Section 3 and with this certificate the company is authorized to affix CE Mark, as shown below, and Notified Body Number on the products in the scope of the examined quality system. Notified Body has the right to carry out required surveillance audits according to Medical Device Directive Annex II Section 5.

For the CE marking of Class III product(s) in the scope of this certificate, EC Design Examination Certificate issued according to Medical Device Directive Annex II Section 4

CE

Certificate Number: 1783 - MDD - 118



SEZAÍ DOĞAN

SEZAİ DOĞAN Director of Directives ANKARA Rev 00, 17/05/2019

This certificate is valid only with the TSE Notified Body Number 1783 seal.



AT TASARIM INCELEME BELGESI

(EC DESIGN EXAMINATION CERTIFICATE)

BELGE NO (CERTIFICATE NO): 1783 - MDD- 119

Aşağıda adı ve adresi yazılı üreticinin tasarımı;

(design of the manufacturer)

MERIL LIFE SCIENCES PVT LTD

MUKTANAND MARG, CHALA, VAPI-396191 GUJARAT INDIA (Merkez Adres/Head Office)

MUKTANAND MARG, CHALA, VAPI-396191 GUJARAT INDIA (Üretim Adresi/Manufacturer Address)

93/42/AT – Tibbi Cihaz Yönetmeliği (EK-II (BÖLÜM 4)) gereklerine göre incelenmiş ve belgelendirilmiştir.

Has been examined and certified according to 93/42/EEC Medical Device Directive (ANNEX II (Article 4))

SİROLİMUS SALINIMLI BİYOEMİLEBİLİR VASKÜLER SCAFFOLD SİSTEMİ SIROLIMUS ELUTING BIORESORBABLE VASCULAR SCAFFOLD SYSTEM (MERES 100^{TM} LINEAGE)

1783

Onaylanmış Kuruluş No Notified Body Number:

Belge Veriliş Tarihi Date of Issue: 17.05.2019

Belge Geçerlilik Tarihi Valid Until: 17.05.2024

Proje Kayıt No Project Registration Number: 1542-18/12817, 1543-18/12823

GMDN Kodu GMDN Code: 56304

Tam Kalite Güvence Belgesi No: 1783-MDD-118

Full Quality Assurance Certificate Number:

Tasarım Dosyası Değerlendirme Rapor No: 1542-MDD-099/2019-02

Design Dossier Review Report Number:

Belge Değişiklik Tarihi / Nedeni:

Date / Reason of the Certificate Revision

Bu belge ekleriyle birlikte geçerlidir. Ekleriyle birlikte 4 sayfadır. (This certificate is valid only with attached annex, if any 4 pages including this page)

AT Tasarım İnceleme Sertifikası, Sınıf III ürünler için, Tam Kalite Güvence (EK-II Bölüm 4 Hariç) belgesinin bir parçasıdır.

For Class III products EC Design-Examination Certificate is part of the Full Quality Assurance Certificate (MDD Annex II Article 4 Excluded)



SEZAİ DOĞAN Direktifler Müdürü Director of Directives Ankara Rev00, 17/05/2019

Bu belge ancak TSE- Onaylanmış Kuruluş Numarası 1783 mührü ile geçerlidir. This certificate is only valid if sealed with "TSE- European Notified Body Number 1783".



AT TASARIM İNCELEME BELGESİ EKİ ANNEX TO THE EC DESIGN EXAMINATION CERTIFICATE BELGE NO (CERTIFICATE NO): 1783 – MDD –119

Ürünün Kullanım Amacı (Intended Use of the Product)

MeRes100[™] Lineage BRS perkütan transluminal koroner anjiyoplasti ve scaffold prosedürlerine elverişli hastaların nativ koroner arterlerinde de novo ve stent için restenotik lezyonlara bağlı semptomatik iskemik hastalık gelişmesi durumda koroner luminal çapın geliştirilmesi için endikedir.

 $MeRes100^{TM}$ Lineage BRS is indicated for improving the coronary luminal diameter in patients with symptomatic ischemic disease due to de novo and in-stent restenotic lesions in native coronary arteries in patients eligible for percutaneous transluminal coronary angioplasty (PTCA) and scaffolding procedure.

Ürün Tipi (Product Type):

MeRes100TM Lineage BRS aşağıdakilerden oluşmaktadır-

- Poli-L-Laktid (PLLA) polimerinden oluşan balonla genişleyebilen scaffold.
- Anit-proliferatif ilaç ve polimer karışımından oluşan scaffold kaplaması:
 - O Anti-proliferatif ilaç Sirolimus (Rapamisin olarak da bilinir)
 - o İlaç rezervuarı ve ilaç salım platformu olarak davranan biyouyumlu, biyobozunur (yardımcı madde) Poli-DL-Lactid (PDLLA) polimer kaplaması.
- Altı çift platinum isaretleyici
- Rapid-exchange scaffold taşiyıcı PTCA balon kateter

MeRes 100TM Lineage BRS comprises of following components-

- A balloon-expandable scaffold made from polymer Poly-L-Lactide (PLLA).
- A scaffold coating that consists of a blend of anti-proliferative drug and polymer:
 - O Anti-proliferative drug Sirolimus (also known as Rapamycin)
 - Bio-compatible, bio-degradable polymer (Excipient) Poly-DL-Lactide (PDLLA) coating which acts as drug reservoir and drug release platform.
- Six pairs of platinum markers
- A rapid-exchange scaffold delivery PTCA balloon catheter

Mevcut Scaffold	Mevcut Scaffold Uzunluğu / Available Scaffold Length (mm)							
Çapı / Available Scaffold Diameter (mm)	13	16	19	24	29	32	37	40
2.25	MRL22513	MRL22516	MRL22519	MRL22524	MRL22529	MRL22532	MRL22537	MRL22540
2.50	MRL25013	MRL25016	MRL25019	MRL25024	MRL25029	MRL25032	MRL25037	MRL25040
2.75	MRL27513	MRL27516	MRL27519	MRL27524	MRL27529	MRL27532	MRL27537	MRL27540
3.00	MRL30013	MRL30016	MRL30019	MRL30024	MRL30029	MRL30032	MRL30037	MRL30040
3.25	MRL32513	MRL32516	MRL32519	MRL32524	MRL32529	MRL32532	MRL32537	MRL32540
3.50	MRL35013	MRL35016	MRL35019	MRL35024	MRL35029	MRL35032	MRL35037	MRL35040
Oles Mande	MRL40013	MRL40016	MRL40019	MRL40024	MRL40029	MRL40032	MRL40037	MRL40040

Marka (Trademark):

MeRes100TM Lineage

Ürünün Tanımı ve İşlevinin Açıklaması (Identification of the Product and description of its functioning)

MeRes100TM Biyoemilebilir Scaffold sistemi balonla genişleyebilen poli-l-laktid (PLLA) biyoemilebilir vasküler scaffold, scaffoldu kaplayan polimer ve sirolimus karışımı kaplaması, scaffold üzerine yerleştirilmiş 6 çift radyopak platinyum işaretleyici ve rapid exchange PTCA balon kataterinden oluşmaktadır. Scaffold balon katater üzerine oturtulmuş ve iki platinyum-iridyum işaretleyici arasına sabitlenmiştir.

PTCA balon kateter, scaffoldu hedef bölgeye taşıyan ve yerleştiren bir taşıyıcı sistem görevi görür. Balon, salin ile kontrast madde karışımı kullanılarak hidrolik basınç uygulaması ile şişirilir. Monte edilmiş scaffold, balonun şişmesi nedeniyle genişler ve hedef bölgeye yerleşir. Balon söndükten sonra stent genişlemiş halde kalır. Taşıyıcı sistem çıkarılır ve sscaffold, hedef bölgeye kalıcı olarak implante edilir; böylece luminal çapta iyileşme ve kan akışının restorasyonu sağlanmış olur. Bu genişleyebilen biyoemilebilir scaffoldlar daralmış arterleri fiziksel olarak destekler ve iskemik arter hastalığının semptomlarını azaltır. Scaffold zaman içerisinde bozunarak vasküler fizyolojinin restorasyonu sağlanır. Scaffoldun üzerine kaplanmış olan ilaç (yani sirolimus), ikili bir etki mekanizmasına sahiptir. Enflamatuar hücre fonksiyonunu modüle eder ve düz kas hücresi proliferasyonunu bloke eder. Sirolimus ilacı scaffold üzerine, ilaç rezervuarı ile ilaç salım platformu görevi gören, biyobozunur özellikte olan ve iyi bilinen bir polimer kombinasyonu ile birlikte kaplanır. Polimer, ilaç ile neredeyse eşzamanlı olarak salınır ve böylece polimere eğilim minimuma indirilmiş olur.

MeRes 100^{TM} Lineage BRS comprises of a balloon expandable bioresorbable PLLA scaffold, a scaffold coating that consists of a blend of sirolimus drug and polymer, six pairs of platinum markers and a rapid exchange Rx balloon dilatation catheter. Scaffold is mounted on a rapid exchange balloon dilatation catheter between two platinum iridium radiopaque markers bands placed on the inner lumen within the balloon segment.

The Rx balloon catheter serves as the scaffold delivery system that delivers and deploys the scaffold to the target site. The balloon is inflated by hydraulic pressurization using mix of saline and contrast medium. The mounted scaffold expands due to inflation of balloon and is deployed to the target site. The scaffold remains expanded after deflation of the balloon. The delivery system is removed and the scaffold is implanted at the target site resulting in improving luminal diameter and restoration of blood flow. These expandable bioresorbable scaffold devices physically support narrowed arteries to alleviate symptoms of ischemic artery disease. The bioresorbable scaffold degrades with time leading to the restoration of vascular physiology. The drug coated on the scaffold i.e. Sirolimus has dual mechanism of action. It modulates inflammatory cell function and blocks smooth muscle cell proliferation. Sirolimus drug is coated on stent with a well-known bio-compatible, biodegradable polymer coating which acts as drug reservoir and drug release platform. The polymer elutes almost simultaneously as the drug, thus minimizing the propensity for a polymer.



AT Tasarım İncelemesine Konu Olan Cihaz Tipi İçin Temin Edilen Teknik Doküman Listesi (List of the technical documentation provided for the appliance type relating to EC Desing Examination)

Teknik Doküman Referansı (Reference of the technical documentation)
Teknik Dosyasında (In Technical File)
Teknik Dosyasında (In Technical File)
Teknik Dosyasında (In Technical File)

Bu belge ancak TSE- Onaylanmış Kuruluş Numarası 1783 mührü ile geçerlidir. This certificate is only valid if sealed with "TSE- European Notified Body Number 1783".





EC Certificate

Directive 93/42/EEC Annex II, excluding Section 4 Full Quality Assurance System Medical Devices

Registration No.: HD 60118775 0001

Report No.: 17054840 003

Manufacturer: Shunmei Medical

Co., Ltd.

R401 of building B, No.8 of 1st Jinglong Road, Baolong

Industrial Zone, LongGang District, 518116 Shenzhen, Guangdong

China

Products: Medical Devices

(see attachment for products and additional sites included)

Notified Body

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Replaces Approval, Registration No.: HD 60107860 0001

Expiry Date: 2021-03-09

The Notified Body hereby declares that the requirements of Annex II, excluding section 4 of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex II, section 5 of the aforementioned directive. For placing on the market of class III devices covered by this certificate an EC design-examination certificate according to Annex II, section 4 is required.

Effective Date: 2017-08-29

Date: 2017-08-29

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg

TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 93/42/EEC concerning medical devices with the identification number 0197.



Doc. 1/2, Rev. 0

TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment to Certificate

Registration No.:

HD 60118775 0001

Report No .:

17054840 003

Manufacturer:

Shunmei Medical

Co., Ltd.

R401 of building B, No.8 of 1st Jinglong Road, Baolong

Industrial Zone, LongGang District, 518116 Shenzhen, Guangdong

China

Products:

- Disposable Pressure Transducers
- Hemodialysis Catheter Kits
- Connecting Tubing
- Introducer Sets
- Guide Wires
- Hemostasis Valve Set
- Ureteral Stent Set
- Introducer Needle
- Angiographic Syringe
- Closed Suction Kit
- Drainage Catheter
- Tracheostomy Tube
- Percutaneous Nephrostomy Sets
- Cervical Ripening Balloon
- Postpartum Balloon with Rapid Instillation Components

Date: 2017-08-29





Doc. 2/2, Rev. 0

TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Attachment to Certificate

Registration No.:

HD 60118775 0001

Report No.:

17054840 003

Manufacturer:

Shunmei Medical

Co., Ltd.

R401 of building B, No.8 of 1st Jinglong Road, Baolong

Industrial Zone, LongGang District, 518116 Shenzhen, Guangdong

China

Aspects of manufacture concerned with securing and maintaining sterile conditions:

- Stopcocks
- Manifolds
- Balloon Inflation Devices
- Dose Control Syringe
- Manifold Kit
- Angio-closure Pad
- TR-Closure Band
- Needle-free Connector

Sites included:

Floor1-floor3 of building C, No.8 of 1st Jinlong Road, Baolong Industrial Zone, LongGang District, 518116 Shenzhen, Guangdong, China

Huizhou branch of Shunmei Medical Co., Ltd Yifa industrial zone, Dushi village, Pingtan town, HuiYang District, HuiZhou City, China

Date: 2017-08-29







Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No.

CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan

Utah 84095 USA

In respect of:

See certificate scope page.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 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: 2008-10-03

Date: 2019-02-08

Expiry Date: 2023-10-02

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Page 1 of 10

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 No: CE 541900

Certificate Scope:

The design, development and manufacture of sterile angiographic, angioplasty and other procedure kits/packs, angiographic catheters, cardiac catheters, vascular catheters, peripheral catheters, guiding catheters, guide wires (coated and uncoated), vascular trocars, introducer needles, angiographic needles, hemodialysis catheters, introducer devices, dilators, transducers, drainage devices, contrast management devices, embolectomy devices, snare devices, hemostasis devices, balloon inflation systems, scalpels, tubing, manifolds/stopcocks, valves, syringes, tracheobronchial stent systems, esophageal stent systems, biliary stent systems, stent positioning system intended for coronary or renal interventional procedures, Peritoneal Dialysis Catheters, accessories and kits, embolization particles, biopsy instruments and accessories, vascular grafts, graft accessory component kits, orthopedic bone cement, bone cement delivery devices/accessories, orthopedic surgical instruments and RF tumor ablation systems for orthopedic applications, percutaneous transluminal angioplasty (PTA) catheters, caps for the disinfection of vascular access connectors, bipolar coagulation probes and all related accessories.

Those aspects of Annex II related to securing and maintaining sterility in the manufacture of angiographic, angioplasty and other procedure kits/packs, anesthesia conduction catheter fixation devices, catheter flush devices, infusion systems, syringes, suture retention devices, torque devices, drainage/waste/sharps collection devices, surgical/general purpose organizers, abdominal binders, labeling sets, compression devices, balloon inflation systems, non-vascular balloon catheter systems and all related accessories.

Those aspects of Annex II related to metrology in the manufacture of syringes, pressure monitors, tracheal measuring devices, balloon inflation systems and all related accessories.

First Issued: **2008-10-03** Date: **2019-02-08** Expiry Date:

Expiry Date: 2023-10-02

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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.





Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095 USA

NBOG code(s)	Device description	Intended purpose	
Class III			
MD 0102/MD 0106; MDS7006	Angiographic and Guide Catheters	See CE 538238	
MD 0102 MDS7006	Drainage Catheters	See CE 541480	
MD 0106, MDS7006	Merit Microcatheters	See CE 553250	
MD 0106, MDS7006	EN Snare Endovascular Snare System	See CE 555846	
MD 0106, MDS7006	InQwire® Diagnostic Guide Wires, InQwire® Amplatz Guide Wires	See CE 560101	
MD 0106, MDS7006	Merit Embolectomy Catheters	See CE 561259	
MD 0106, MDS7006	Ostial Pro Stent Positioning System	See CE 585005	
MD 0106, MDS7006	ONE Snare Endovascular Snare System, ONE Snare Endovascular Microsnare System	See CE 590890	
MD 0102, MDS7006	Hemodialysis Catheters	See CE 606106	
MD 0106, MDS7006	Merit SureCross™ Support Catheter	See CE 612029	
MD 0102, MDS7006	HeRO Graft	See CE 650631	
MD 0106, MDS7006	SwiftNINJA Steerable Microcatheters	See CE 667696	
MD 0106, MDS7006	True Form Reshapable Guide Wire	See CE 669204	

First Issued: 2008-10-03 Date: 2019-02-08 Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095

USA

NBOG code(s)	Device description	Intended purpose
Class IIb		
MD 0200, MDS7006	Biliary Catheter (RBC), Biliary Drainage Cath. (RBDC), CirQ Nephrostomy Catheter	intended for drainage of bile within the biliary system
MD 0200, MDS7006	ReSolve Locking Catheter (RLC)	intended for percutaneous drainage of fluids from body cavities for up to 90 days.
MD 0106, MDS7006	ALIMAXX-ES, EndoMAXX, EndoMAXX EVT	used in the treatment of malignant neoplasms for the purpose of palliating the airway.
MD 0106, MDS7006	AERO Tracheobronchial Stent, AeroMINI, AERO Delivery System	indicatedfor the use in the treatment of tracheobronchial strictures and airway compressions (stenosis) produced by malignant neoplasms
MD 0106, MDS7006	Biliary Stents & Delivery System: ALIMAXX-B	indicated for the palliation of malignant strictures in the biliary tree
MD 1104, MDS7006	Bipolar Coagulation Probe and related accessories	Probes function as conventional electro- coagulation devices when supplied withcurrent from a standard bipolar electro-surgical generator. The probes are intended to provide hemostasis throughout the gastrointestinal tract.
MD 0102, MDS7006	Flex-Neck® Classic, Infant, ARC, ExxTend Catheters	intended for implantation for more than 30 days to carry fluid into and out of the abdomen

First Issued: 2008-10-03 Date: 2019-02-08 Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To: Merit Medical Systems, Inc.

1600 West Merit Parkway South Jordan

Utah

84095 USA

NBOG code(s)	Device description	Intended purpose
Class IIb		
MD 0106, MDS7006	Peritoneal Dialysis Catheter Embedding Tool	indicated for embedding the external portion of most PD catheters subcutaneously in anticipation of future retrieval of the part of the catheter
MD 0202, MDS7006	StabiliT ER2 Bone Cement and Saturate Mixing System	intended for use in treatment of pathological fractures of the vertebrae using vertebroplasty or kyphoplasty procedure
MD 1402, MDS7006	SpineSTAR Tumor Ablation Systems (instruments and kits)	intended for the ablation of tumor within the vertebral body. It heats targeted tissue in contact with the electrode
MD 1402 (non-sterile)	MetaSTAR RF Generator	intended to generate and control the delivery of RF energy for palliative treatment in spinal procedures by ablation of metastatic malignant lesions in a vertebral body
MD 0200, MDS7006	BioSphere Bearing nsPVA	used for the embolization of peripheral hypervascularized tumors, including leiomyoma uteri and peripheral arteriovenous malformations (AVMs)
MD 0202, MDS7006	StabiliT Vertebral Augmentation & Vertebroplasty Kits (Class IIb kits under article 12)	for the treatment of pathological fractures of the vertebrae using a vertebroplasty or kyphoplasty procedure

First Issued: 2008-10-03 Date: 2019-02-08 Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095 USA

NBOG code(s)	Device description	Intended purpose
Class IIb		
MD 0202, MDS7006	StabiliT Vertebral Augmentation & Vertebroplasty Kits (Class IIb kits under article 11)	for the treatment of pathological fractures of the vertebrae using a vertebroplasty or kyphoplasty procedure
Class IIa		
MD 0102, MDS7006	Fountain & Mistique Infusion Catheters	NA for class IIa devices
MD 0102, MDS7006	One Step Centesis Drainage Catheter	NA for class IIa devices
MD 0102, MDS7006	ReSolve Non-Locking Catheter (RNL), Resolve Dilator	NA for class IIa devices
MD 0104, MDS7006	Intelliflator & Merit Monitor (IntelliSystem), DiamondTOUCH, Monarch, Blue Diamond, Endotek Digital Inflation Syringes	NA for class IIa devices
MD 0106, MDS7006	Introducer, Mini Access, Radial, Plastic Jacket Guide Wires	NA for class IIa devices
MD 0106, MDS7006	MAXXWIRE & ENDOWIRE (Aero) Guide Wires	NA for class IIa devices
MD 0106, MDS7006	Manifolds, Stopcocks, Rotating Adapters, Flow Switch, TRAM	NA for class IIa devices
MD 0106, MDS7006	MAK/SMAK, MAK-NV, Vessel Dilator	NA for class IIa devices
MD 0106, MDS7006	Valve Adapter	NA for class IIa devices
MD 0102, MDS7006	High Pressure Contract Injection and Pressure Monitoring Tubing	NA for class IIa devices
MD 0104, MDS7006	MeriTrans™ and Argotrans Disposable Transducers	NA for class IIa devices

First Issued: 2008-10-03 Date: 2019-02-08 Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095

USA

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NBOG code(s)	Device description	Intended purpose
Class #6		
MD 0106, MDS7006	Merit Angioplasty Pack (MAP Kits)	NA for class IIa devices
MD 0106, MDS7006	FLO50, PhD, Passage, Access-9, AccessPLUS, DoublePlay, MBA, MBA Plus, Honor, FLO30, FLO40, FLO40XR Hemostasis Valves	NA for class IIa devices
MD 0106, MDS7006	Prelude (PSI & PRO), HVA, PreludeEASE, Prelude short Sheath, Dilator & Obturator	NA for class lla devices
MD 0106, MDS7006	Advance™ Merit Angiographic Needles	NA for class IIa devices
MD 0106, MDS7006	Futura™ Safety Scalpel	NA for class IIa devices
MD 0106, MDS7006	OuTake Catheter Extractor Device	NA for class IIa devices
MD 0102, MDS7006	Contrast Management Systems (high pressure use)	NA for class IIa devices
MD 0102, MDS7006	CT Transfer Set - FAS	NA for class IIa devices
MD 0106, MDS7006	Flow Guard™ Valved Peelable Introducer	NA for class IIa devices
MD 0106, MDS7006	Hart Chiba & Trocar Style Needles	NA for class IIa devices
MD 0106, MDS7006	Merit Advance Angiographic Safety Needles	NA for class IIa devices
MD 0106, MDS7006	Laureate Guide Wire (peripheral and urological)	NA for class IIa devices
MD 0106, MDS7006	Peritoneal Dialysis Implantation Kits	NA for class IIa devices
MD 0106, MDS7006	Corvocet Coaxial Introducer and Biopsy System	NA for class IIa devices
MD 0106, MDS7006	HeRO Accessory Kit	NA for class IIa devices
MD 0106, MDS7006	Hydraulic Assembly and Hydraulic Master Syringe Assembly	NA for class IIa devices
MD 1402, MDS7006	Activation Element	NA for class IIa devices

First Issued: **2008-10-03** Date: **2019-02-08** Expiry Date: **2023-10-02**

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan

Utah 84095 USA

110

NBOG code(s)	Device description	Intended purpose
Class Ith		
MD 1402 (non-sterile)	Multiplex Controllers	NA for class IIa devices
MD 0103, MDS7006	StabiliT and STAR Instruments	NA for class IIa devices
MD 0106, MDS7006	Advocate PTA Balloon Catheter	NA for class IIa devices
MD 0108, MDS7006	DualCap System	NA for class IIa devices
MD 0106, MDS7006	Osseoflex Access Cannulas and Stylets	NA for class IIa devices
MD 0106, MDS7006	Osseoflex Steerable Needles	NA for class IIa devices
MD 0106, MDS7006	Bone Filler Devices	NA for class IIa devices
MD 0106, MDS7006	Bone Drills	NA for class IIa devices
MD 0106, MDS7006	Pursue (Hi-Lex) Microcatheter	NA for class IIa devices
MD 0106, MDS7006	Osseoflex Cement Delivery System	NA for class IIa devices
MD 0106, MDS7006	Bone Marrow Aspiration Needle	NA for class IIa devices
Class Is		
MDS7006	Squirt Fluid Dispensing System	NA for class Is devices
MDS7006	External Vascular Compressors / RadStat Radial Artery Compression System; Compression Discs	NA for class Is devices
MDS7006	Tags / PAL Pen and Labels; Custom PAL Labels; PAL Pen	NA for class Is devices

First Issued: 2008-10-03 Date: 2019-02-08 Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095 USA

NBOG code(s)	Device description	Intended purpose
Class Is		
MDS7006	Basins / Backstop® Disposal Basin; Ministop Disposal Basins; BackStopPlus Disposal Basis; MiniStopPlus Disposal Basin; Triple Play™ Disposal Basin; Dugout® DisposalBasin; RingMaster Guide Wire Basin	NA for class Is devices
MDS7006	Valves / Check Relief Valve; In - Line Check Relief Valve [CRV] Caotiva® Blood Containment Device	NA for class Is devices
MDS7006	Catheter Tubeholders / Revolution Catheter Securement Device	NA for class Is devices
MDS7006	Fluid Administration Set; Fluid Administration Spike; Fluid Management Tube	NA for class Is devices
MDS7006	Merit Miser Contrast Management System	NA for class Is devices
MDS7006	Continuous Flush Devices	NA for class Is devices
MDS7006	Merit Angioplasty Packs	NA for class Is devices
MDS7006	Waste Disposal System	NA for class Is devices
MDS7006	Merit Disposal Depot	NA for class Is devices
MDS7006	Shortstop and Shortstop Advantage Temporary Sharps Holders	NA for class Is devices
MDS7006	Procedure Packs (Custom Kits); Rapid Response Kits	NA for class Is devices
MDS7006	Connection Tubes	NA for class Is devices
MDS7006	Syringes	NA for class Is devices

First Issued: 2008-10-03

Date: 2019-02-08

Expiry Date: 2023-10-02

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Supplementary Information to CE 541900

Issued To:

Merit Medical Systems, Inc. 1600 West Merit Parkway South Jordan Utah 84095 USA

NBOG code(s)	Device description	Intended purpose
Class Is		
MDS7006	Syringes	NA for class Is devices
MDS7006	Merit Drainage Depot	NA for class Is devices
MDS7006	Basix™ Inflation Syringe; BasixCOMPAK Inflation Syringe; BasixTOUCH Inflation Syringe	NA for class Is devices
MDS7006	Angiography and Angioplasty Procedures Accessories of Class I Sterile / RXP® Rapid Exchange Prep Syringe; Guide Wire Insertion Tool; Pin Vice Torque Device; H20 TORQ Device; SeaDragon Torque Device	NA for class Is devices
Class Im		
MD 1301 (non-sterile)	Monitors / IntelliSystem® II Color Monitor	NA for class Im devices
MD 0104 (non-sterile)	AeroSIZER Stent Measuring System	NA for class Im devices

First Issued: **2008-10-03**

Date: 2019-02-08

Expiry Date: 2023-10-02

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EC CERTIFICATE

Number: 3812454CE01

Full Quality Assurance System

Directive 93/42/EEC on Medical devices, Annex II excluding (4)

(Devices in Class IIa, IIb or III and Devices in Class I in sterile conditions and sterilised systems or procedure packs)

Manufacturer:

Boston Scientific Corporation

300 Boston Scientific Way Marlborough, MA 01752 United States Of America

For the product category(ies)

Medical Devices and Accessories for Minimally Invasive biliary, cardiovascular (including cardiovascular interventions), electrosurgical, endoscopic surgical, endoscopy, gastroenterology, gynaecology, nephrology, neurology (including neurovascular), peripheral (including peripheral interventions) and urology procedures.

DEKRA grants the right to use the EC Notified Body Identification Number illustrated below to accompany the CE Marking of Conformity on the products concerned conforming to the required Technical Documentation and meeting the provisions of the EC-Directive which apply to them:

0344

Documents, that form the basis of this certificate:

Certification Notice 3812454CN, initially dated 1 July 2014 Addendum, initially dated 1 July 2014

DEKRA hereby declares that the above mentioned manufacturer fulfils the relevant provisions of /Besluit Medische Hulpmiddelen', the Dutch transposition of the Council Directive 93/42/EEC of June /14, 1993 concerning Medical devices, including all subsequent amendments. The manufacturer has implemented a quality assurance system for design, manufacture and final inspection, that covers the aspects of manufacture concerned with securing and maintaining sterile conditions, for the above mentioned product category in accordance to the provisions of Annex II Council Directive 93/42/EEC of June 14, 1993 and is subject to periodical surveillance. Additionally, DEKRA hereby declares that the manufacturer fulfils the relevant provisions as specified in Annex I of Commission Regulation 722/2012 of 8 August, 2012 concerning medical devices manufactured utilising tissue of animal origin. For placing on the market of Class III devices an additional EC design examination certificate according to Annex II (4) is mandatory. The necessary information related to the quality management system of the manufacturer, including facilities and the reference to the relevant documentation, of the products concerned and the assessments performed, are stated in the Certification Notice which forms an integrative part of this certificate.

This certificate is valid until: 14 December 2023 Revised: 21 December 2018 Issued for the first time: 1 July 2014 Reissued: 14 December 2018

DEKRA Certification B.V.

B.T.M. Holtus Managing Director J.A. van Vugt Certification Manager

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DEKRA Certification B.V. is Notified Body with ID no 0344

ADDENDUM

Belonging to certificate: 3812454CE01

CE MARKING OF CONFORMITY MEDICAL DEVICES

Medical Devices and Accessories for Minimally Invasive biliary, cardiovascular (including cardiovascular interventions), electrosurgical, endoscopic surgical, endoscopy, gastroenterology, gynaecology, nephrology, neurology (including neurovascular), peripheral (including peripheral interventions) and urology procedures.

Issued to:

Boston Scientific Corporation

300 Boston Scientific Way Marlborough, MA 01752 United States Of America

This certificate covers the following location(s):

Location Code	Company name / address	Location Code	Company name / address
MAR2	Boston Scientific Corporation 300 Boston Scientific Way Marlborough, MA 01752 USA	CR2	Boston Scientific Corporation 2546 First Street, Propark El Coyol Alajuela Costa Rica
COR	Boston Scientific Limited Business & Technology Park Model Farm Rd Cork, Ireland	GAL	Boston/Scientific/Limited Ballybrit Business/Park Galway, Ireland
COV	Boston Scientific Corporation 8 Industrial Drive Coventry, RI 02816 USA	KER	Boston Scientific International BV European Centre of Operations Vestastraat 6, 6468 EX Kerkrade, The Netherlands

DEKRA Certification B.V.

B.T.M. Holtus Managing Director

J.A. van Vugt Certification Manager

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Issued to:

Boston Scientific Corporation

300 Boston Scientific Way Marlborough, MA 01752 United States Of America

This certificate covers the following location(s):

Location Code	Company name / address	Location Code	Company name / address
CR1	Boston Scientific Corporation 302 Parkway Global Park, Heredia Costa Rica	MAR	Boston Scientific Corporation /100 Boston/Scientific Way /Marlborough, MA 01752 /USA
MG2	Boston Scientific Corporation Two Scimed Place Maple Grove, MN 55311 USA	(SJ2/	Boston Scientific Corporation 150 Baytech Drive San Jose, CA 95134 USA
PL2	Boston Scientific Corporation 5905 Nathan Lane Plymouth, MN 55442 USA	SPE	Boston Scientific Corporation 780 Brookside Drive Spencer, IN 47460 USA

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B.T.M. Holtus Managing Director

J.A. van Vugt Certification Manager

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Issued to:

Boston Scientific Corporation

300 Boston Scientific Way Marlborough, MA 01752 United States Of America

This certificate covers the following location(s):

QUI Boston Scientific Corporation Marina Bay Customer Fulfillment Center 500 Commander Shea Blvd Quincy, MA 02171 USA

Initial date: 1 July 2014

Revision date: 21 December 2018

DEKRA Certification B.V.

B.T.M. Holtus Managing Director J.A. van Vugt Certification Manager

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