Către Agenția Medicamentului și Dispozitivelor Medicale

NOTIFICARE

pentru înregistrarea dispozitivelor medicale în Registrul de stat al dispozitivelor medicale

nr. 3 din 13.10.2023

Solicitantul <u>Oxivit Med SRL</u>, cu sediul <u>mun.Chisinau</u>, <u>MD-2020</u>, <u>Stradela Studentilor</u>, <u>6b</u>, tel./fax: <u>+37368781333</u>_, e-mail <u>oxivit.medical@gmail.com</u>, solicit înregistrarea în Registrul de stat al dispozitivelor medicale a următoarelor categorii și tipuri de dispozitive medicale pentru introducerea și punerea la dispoziție pe piață a:

- Telescope Guide Extension Catheter: TELE6F, TELE7F

Se anexează următoarele acte:

<u>Declarație pe proprie răspundere</u>

<u>CE certificate</u>

<u>Declaratie de conformitate</u>

<u>Scrisoare de imputernicire</u>

ata	13/10/2023	Semnatur

Tabelul de recepționare a notificării

(se completează de către Agenție în momentul depunerii notificării de către solicitant)

Comentarii cu privire la acceptul/refuzul	
recepționării notificării, inclusiv motivul	
refuzului	
Data/nr. de ordine atribuit notificării de către	
Agenție (în cazul acceptării recepționării)	
Numele, prenumele, funcția persoanei	
responsabile de recepționarea dosarului	
Semnătura persoanei responsabile	

Către Agenția Medicamentului și Dispozitive Medicale

DECLARAȚIE PE PROPRIE RĂSPUNDERE

Solicitant: <u>Oxivit Med SRL</u> , cu sediul	mun.Chisinau, MD-
2020, Stradela Studentilor, 6b,	
declar pe proprie răspundere, cunoscând preveder Republicii Moldova cu privire la falsul în declarații, că do	
pentru notificarea dispozitivului medical:	ocumentele și ducele furnizace
 Telescope Guide Extension Catheter: TELE6F, TE Sunt autentice și corespund realității. 	LE7F
, .	
Numele, prenumele și funcția	Semnătura
Kojevnikov Dmitrii, director	
	Data 13/10/2023

Medtronic META FZ-LLC

Injaz Building, 3rd floor

Dubai Knowledge Park, P.O. Box 500638

Dubai, United Arab Emirates

www.medtronic.com
Tel: +971 4 818 2666

TO WHOM IT MAY CONCERN

Date: 29.05.2023

We hereby certify that, pursuant to a non-exclusive distribution agreement expiring on 31 May 2024 between Medtronic META FZ-LLC (the "Company"), a Medtronic company organized under the laws of the United Arab Emirates having its principal place of business at 3rd Floor, Injaz Building, Dubai Knowledge Park, Dubai, United Arab Emirates, and OXIVIT-MED, SRL, a company having its principal place of business at Bld. Moscova 14/1., Chisinau, MD-2068, Republic of Moldova, the latter is authorized to act as the Company's distributor in Republic of Moldova for the below listed product lines:

- Cardiac Ablation Solutions;
- Cardiac Rhythm Management.
- Cardiovascular Diagnostics & Services;
- Cardiac Surgery
- Coronary & Renal Denervation
- Structural Heart & Aortic
- Peripheral & Endovenous
- Cranial & Spinal Technologies
- Neuromodulation
- Pelvic Health
- Ear, Nose & Throat
- Neurovascular
- Surgical Innovation
- Gastrointestinal
- Patient Monitoring
- Respiratory interventions
- Diabetes

This letter is valid until 31 May 2024 and can also be used with the Competent Authorities in Moldova for notification of medical devices manufactured by attached Medtronic and Covidien Legal Manufacturers, or other regulatory purposes.

Medtronic META FZ-LLC

M. N. AL-A NF

Name: Muzahim Al Ani Title: Authorized Signatory

APPENDIX 1

Medtronic Manufacturing Facilities/Legal Manufacturers:

- 1. Medtronic, Inc., 710 Medtronic Parkway, Minneapolis MN 55432, USA
- 2. Medtronic, Inc., 7000 Central Avenue N.E., Minneapolis MN 55432, USA
- 3. Medtronic, Inc., 3800 Annapolis Lane, Minneapolis MN 55447, USA
- 4. Medtronic, Inc., 8200 Coral See Street N.E., Mounds View, MN 5512, USA
- 5. Medtronic Europe Sàrl, Route du Molliau 31, Case Postale, CH-1131 Tolochenaz, Switzerland
- 6. Medtronic Bakken Research Center B.V., Endepolsdomein 5, 6229 GW Maastricht, The Netherlands
- 7. Medtronic CoreValve LLC, 1851 E. Deere Avenue, Santa Ana, CA 92705, USA
- 8. Medtronic MiniMed, 18000 Devonshire street, Northridge CA 91325-1219, USA
- 9. Medtronic Heart Valves Division, 1851 East Deere Avenue, Santa Ana, CA 92705, USA
- 10. Medtronic Heart Valves, 1941 Blair Avenue, Santa Ana CA 92705, USA
- 11. Medtronic Fabrication S.A.S. Zone Industrielle SUD, Route D'Anor, 59610 Fourmies, France
- 12. Medtronic Sofamor Danek USA, Inc., 1800 Pyramid Place, Memphis, TN 38132, USA
- 13. Medtronic Sofamor Danek USA, Inc., 4340 Swinnea Road, Memphis TN 38118, USA
- 14. Medtronic Sofamor Danek Manufacturing, 2500 Silveus Crossing, Warsaw IN 46582, USA
- 15. Medtronic Sofamor Danek Deggendorf GmbH, Werfstrasse 17, 94469 Deggendorf, Germany
- 16. Medtronic Sofamor Danek Deggendorf GmbH, Ulrichsberger Str. 17, 94469 Degendorf, Germany
- 17. Medtronic Xomed, Inc. 6743 Southpoint Drive North, Jacksonville FL 32216, USA
- 18. Medtronic Xomed Instrumentation S.A.S., Le Pavillon, Saint Aubin Le Monial, 03160 Bourbon-L'Archambault, France
- 19. Medtronic Xomed, Inc., 950 Flanders Road, Mystic CT 06355, USA
- 20. Medtronic Xomed, Inc., 80 Davids Drive # 5, Hauppauge, NY 11788, USA
- 21. Medtronic Powered Surgical Solutions, 4620 North Beach Street, Forth Worth, Texas 76137, USA
- 22. Medtronic Navigation, Inc. (Littleton), 300 Foster Street, Littleton, MA 01460, USA
- 23. Medtronic Navigation, Inc., 826 Coal Creek Circle, Louisville, CO 80027, USA
- 24. Medtronic Navigation Israel, Ltd. Kochav Yokneam Building, P.O.Box 548, 20692 Yokneam Elit, Israel
- 25. Kyphon Sàrl, Pierre-à-Bót 97, 2000 Neuchâtel, Switzerland
- 26. AF Solutions, Medtronic Inc., 8200 Coral Sea Street, Mounds View, Minneapolis MN 55112, USA
- 27. Medtronic Ireland, Parkmore Business Park West Galway, Ireland
- 28. Medtronic Vascular, 35-37a Cherry Hill Drive, Danvers, MA 01923, USA
- 29. Medtronic Vascular, 3576 Unocal Place, Santa Rosa, CA 95403, USA
- 30. Medtronic Puerto Rico Operations Co., Juncos, Road 31, Km. 24, Hm 4, Ceiba Norte Industrial Park, Juncos PR 00777, USA
- 31. Medtronic Puerto Rico Operations Co., Villalba, Rd 149, Km 56.3, Call Box 6001, PR 00766 Villalba, USA
- 32. Medtronic Puerto Rico Operations, Co., Road 909 Km. 0.4 Bo, 00792 Barrio Mariana, Humacao, Puerto Rico, USA
- 33. Medtronic Puerto Rico Operations, Co., Parcela ≠21, Catano Industrial Park, Dr. John A. Smith Street, 00792 Humacao, Puerto Rico, USA
- 34. Medtronic Mexico S. de R.L. de CV, Av. Paseo Cucapah, 10510 El Lago, C.P. 22210 Tijuana, Baja California, Mexico
- 35. Medtronic Mexico EG, Carreta Internacional Guadalajara Nogales, Empalme, Sonora, Mexico 85340
- 36. Medtronic Singapore Operations Pte. Ltd., 49 Changi South Avenue 2, Nasaco Tec Centre, 486056 Singapore
- 37. Medtronic Atrial Fibrillation Technologies (AFT), 8200 Coral Sea Street NE, Mounds View, Minneapolis MN 55112, USA
- 38. Medtronic Perfusion Systems, 7611 Northland Drive, Minneapolis, MN 55428, USA
- 39. Medtronic Perfusion Systems, 18501 East Plaza Drive, Parker C0 80134-9061, USA
- 40. Medtronic Cardiac Surgery Division Europe, Valkenhuizerlaan 16, 6466 ND Kerkrade, The Netherlands

- 41. Medtronic Neuromodulation, 800 53rd Avenue N.E., Minneapolis MN 55421, USA
- 42. Medtronic Neuromodulation, 7000 Central Avenue N.E. Minneapolis MN 55432, USA
- 43. Invatec S.p.A., 7, Via Martiri Della Libertà, 25030 Roncadelle, Brescia, ItalyInvatec S.p.A., 26-28 Via Industria, 25030 Torbole-Casaglia, Brescia, Italy
- 44. Medtronic Ardian LLC, 1380 Shorebird Way, Mountain View, CA 94043, USA
- 45. Medtronic Neurosurgery, 125 Cremona Drive, Goleta, CA 93117, USA
- 46. Medtronic Advanced Energy, LLC, 180 International Drive, Portsmouth NH 03801, USA
- 47. Vitatron Holding B.V. Endepolsdomein 5, 6229 GW Maastricht, The Netherlands
- 48. HeartWare, Inc.14400 NW 60th Avenue, Miami Lakes, Florida 33014, USA
- 49. BELLCO SOCIETÀ, Unipersonale A r.l., Via Camurana 1, 41037 Mirandola (MO), IT Italy
- 50. Medtronic CryoCath LP., 9000 Autoroute Transcanadienne, Pointe Claire, Quebec H9R 5Z8, Canada
- 51. CardioInsight Technologies Inc., 11000 Cedar Ave, Suite 210, Cleveland, OH 44106
- 52. Medtronic Vascular Inc., 271 Gibraltar Drive, Sunnyvale, CA 94089
- 53. Teleflex Medical, Annacotty Business Park, Annacotty, Co. Limerick, Ireland
- 54. TYRX Inc., 1 Deer Park Dr. Suite G Monmouth Junction, NJ 08852
- 55. HeartWare Inc., 500 Old Connecticut Path Framingham, MA 01701
- 56. Medtronic Sofamor Danek Inc., Airport Logistics Center, Building "A", 4340 Swinnea Road, Memphis, TN 38118, USA
- 57. Medtronic Monitoring Inc., 1410 Energy Park Dr. Saint Paul, MN 55108
- 58. Medtronic Vascular, 3850 Brickway Blvd, Santa Rosa, CA 95403 USA
- 59. Osteotech, Inc., 201 Industrial Way, Eatontown, New Jersey, 07724, USA
- 60. Medtronic Spine LLC, 1860 Barber Ln, Milpitas, CA 95035
- 61. Changzhou Kanghui Medical Inovation Co., Ltd., No. 11, North Chanjiang Road, Xinbei District, Changzhou 213022, P.R. China

List of EC Representatives:

- 1. Medtronic B.V., Earl Bakkenstraat 10, 6422 PJ Heerlen, The Netherlands
- 2. Medtronic Ireland, Parkmore Business Park West, Galway, Irelandkang
- 3. Invatec S.p.A., 7, Via Martiri Della Libertà, 25030 Roncadelle, Brescia, Italy

List of Medtronic Distribution Centers:

- 1. Medtronic B.V., Earl Bakkenstraat 10, 6422 PJ Heerlen, The Netherlands
- 2. Medtronic, Inc. Mounds View Distribution, 2292 Wooddale Dr, Ramsey, 55112 Mounds View, USA
- 3. Medtronic, Inc. Grand Rapids Distribution, 2925 Walkent Ct NW, Walker, MI 49544, USA
- 4. Medtronic, Inc. Blood Management Business, 18501 East Plaza Drive, Parker C0 80134-9061, USA
- 5. Medtronic, Inc. Western Distribution Center, 2385 Railroad Street, Corona, CA 92880, USA
- 6. Medtronic Xomed, Inc. Jacksonville Distribution, 6743 Southpoint Drive North, Jacksonville, FL 32216, USA
- 7. Xomed MicroFrance Distribution, Le Pavillon, Saint Aubin Le Monial, 03160 Bourbon-L'Archambault, France
- 8. Medtronic Sofamor Danek, Inc., Airport Logistics Center, Building "B", 4340 Swinnea Road, Memphis, TN 38118, USA
- 9. Medtronic Powered Surgical Solutions, 4620 North Beach Street, Forth Worth, Texas 76137, USA
- 10. Medtronic Leipzig Distribution 3PL, Dingolfinger Strasse 19, 04349 Leipzig, Germany
- 11. Puerto Rico Distribution Center, Santander Tower, B-7 Tabonuco Street, Suite 1501, Guaynabo PR 00968-3028, USA
- 12. Medtronic, Inc. New York Distribution Center East, Exel 3PL, 699 Kapkowski Road, Suite 300, Elizabeth NJ 07201-2122, USA
- 13. International Shipping Associate, UPS-SCS / Medtronic, 1860 Outerloop Road, Louisville, KY 40219, USA
- 14. Medtronic Navigation, Inc., 826 Coal Creek Circle, Louisville, CO 80027, USA
- 15. Medtronic Neurosurgery, 125 Cremona Drive, Goleta, CA 93117, USA
- 16. Swedesboro, NJ Distribution Center, 1130 Commerce Boulevard, Swedesboro NJ 08085
- 17. Guaynabo Distribution Center Sector La Muda Rd #1, Km 21.1, Guaynabo PR 00971, USA

18. MDT Logistics LLC DC East, 1130 Commerce Blvd Ste 100, Swedesboro, NJ 08085-1765 USA

Covidien Manufacturing Facilities/Legal Manufacturers, EC Representatives and Distribution Centers.

Covidien Ilc (USA)

- 15 Hampshire Street, Mansfield, MA 02048 USA
- 5439 State Route 40 Argyle, NY 12809 USA
- 1430 Marvin Griffin Road Augusta, GA 30906 USA
- 815 Tek Drive Crystal Lake, IL 60039 USA
- 525 North Emerald Road Greenwood, SC 29646 USA
- 1448 Blue Ridge Boulevard Seneca, SC 29672 USA
- 1313 West Grant Boulevard Wabasha, MN 55981USA
- Two Ludlow Park Drive Chicopee, MA 01022 USA
- 400 Maple Street Commerce, TX 75428 USA
- 2010 East International Speedway Boulevard Deland, FL 32724 USA
- 1222 Sherwood Road Norfolk, NE 68701USA
- 150 Glover Avenue, Norwalk, CT 06856 USA
- 60 Middletown Ave, North Haven, CT 06473 USA
- Building 911-67, Sabanetas Industrial Park, Ponce, Puerto Rico 00731
- At 101a Fist Avenue, Waltham, MA 02451, USA
- 5920 Longbow Dr., Boulder, CO 80301, USA
- 15 Crosby Drive, Bedford, MA 10730, USA
- 6135 Gunbarrel Ave., Boulder, CO 80301, USA
- 675 McDonnell Blvd., Saint Louis, MO 63134, USA
- formerly Barrx Medical Inc., 540 Oakmead Parkway Sunnyvale, CA 94085, USA

Covidien Ilc (Canada)

Ludlow Technical products Canada Ltd., 215 Herbert Street Gananoque, Ontario K7G 2Y7 Canada

Covidien Ilc (Mexico)

- 37 Boulevard Insurgentes, Libriamento a la P, La Mesa Tijuana, B.C., Mexico
- Calle 9 Sur No. 125 Cuidad Industrial Tijuana, Mexico CP 22500
- Boulevard Insurgentes 19030 Libramiento, 22225 Tijuana, B.C., Mexico
- Avenue Henequen No. 1181, Parque Industrial Salvarcar, 32573 Ciudad Juarez, Chihuahua, Mexico

Covidien Ilc (Dominican Republic)

Zona Franca de San Isidro, Carretera San Isidro Km 17 Santo Domingo, Dominican Republic

Covidien Ilc (Europe)

- Wilson Way Pool Industrial Estate Redruth, Cornwall TR15 3QN United Kingdom Unit 1, Astley Lane Industrial Estate Swillington Leeds West Yorkshire LS26 8XT, United Kingdom
- IDA Business and Tehcnology Park, Srah Industrial Estate Tullamore, Co. Offaly, Republic of Ireland
- Cornamaddy Athlone, Co. Westmeath, Ireland
- Michael Collins Road Mervue, Co. Galway, Ireland
- Via G. Bove, 2/4/6/8 Mirandola (MO) Italy 41037
- Quedlinburger Strasse 39A Halberstadt, D-38820 Germany

Covidien Ilc (Thailand)

117 Moo 2, Petchkasem Rd. Klongmai Sampran, Nakornpathom, 73110, Thailand

Covidien Ilc (Japan)

SBS Hills 1, 4-10-2 Yoga Setagaya-ku Tokyo 158-8615, Japan

Covidien Ilc Newton (Beacon)

2000 Commonwealth Avenue, 1st Floor, Newton, MA 0246, USA

Covidien AG

Victor Von Bruns-Strasse 19, 8212 Neuhausen am Rheinfall, Switzerland

Covidien Logistics BVBA

Weg naar Zwartberg 239, 3660 Opglabbeek, Belgium

Covidien Ireland Commercial Limited

3rd Floor, Kilmore House, Park Lane, Spencer Dock, Dublin 1

Covidien Medical Products (Shanghai) Manufacturing LLC

Bldg. 10, No. 789 Puxing Rd. Shanghai, P.R. China

Covidien (China) Medical Devices Technology Co., Ltd.

Floor 6, Building 3, #2388 Chenhang Road, 201114 Shanghai, P.R. China

Covidien Deutschland Manufacturing GmbH

Gewerbepark 1, 93333 Neustadt and der Donau, Germany

Covidien Manufacturing Solutions SA

Edificio B20 Calle #2 Zona Franca Coyol Alajuela, Costa Rica

Covidien, formerly known as Valleylab, A Division of Tyco Healthcare Group LP

5920 Longbow Drive Boulder, CO 80301 USA

Covidien, formerly Kendall-Gammatron Co. Ltd.

117 Moo 2, Petchkasem Rd. Klongmai Sampran, Nakornpathom, 73110, Thailand

Covidien, formerly MMJ S.A. de C.V.

Avenue Henequen No. 1181, Parque Industrial Salvarcar, 32573 Ciudad Juarez, Chihuahua, Mexico

EV3 Inc.

4600 Nathan Lane North Plymouth, MN 55442 USA

Micro Therapeutics Inc. d/b/a ev3 Neurovascular

9775 Toledo Way, Irvine, CA 92618 USA

superDimension Inc.

161 Cheshire Lane, Suite 100, Plymouth, Minnesota 55441, USA

Sofradim Production

116 Avenue du Formans, 01600 Trévoux, France

VNUS Medical Technologies, Inc.

5799 Fontanoso Way, San Jose CA 95138, USA

Given Imaging, Inc.

15 Hampshire Street, Mansfield MA 02048, USA

Given Imaging, Itd

2 Hacarmel St. New Industrial Park POB 258 Yoqneam, Ha Zafon 20692 Israel

Given Imaging (Los Angeles) LLC

Los Angeles 5860 Uplander Way Culver City, CA 90230 USA

Given Imaging Vietnam Co. Ltd.

Suite 6A, 6th floor, Standard Factory Buliding, 14th Street, Tan Thuan EPZ, Tan Thuan Dong Ward, District 7, Ho Chi Minh City, Vietnam

Wem Equipamentos Eletrônicos Ltda.

Rua. Marechal Mascarenhas de Morais 550, Parque Industrial Lagoinha, Ribeirão Preto, São Paulo - CEP 14095-120, Brazil

<u>List of OEM - Original Equipment Manufacturers</u>

and Sub-Contracting Facilities:

MindFrame, Inc.

12 Goodyear, Suite 125, Irvine, CA 92618, USA

Buffalo Filter, LLC

5900 Genesee St., Lancaster, NY 14086, USA

SCHÖLLY FIBEROPTIC GmbH

Robert-Bosch-Strasse 1-3, 79211 Denzlingen, Germany

W.O.M. World Of Medicine GmbH

Salzufer 8, 10587 Berlin, Germany

Liebel-Flarsheim Company Llc

2111 East Galbraith Road, Cincinnati, OH 45237 USA

Mediquip SDN. BHD.

- Padang Lati, Mukim Paya, P.O. Box 25, 01700 Kangar, Perlis, Malaysia
- Batu 5, Padang Lati, Jln. Santan, Kangar, 02450 Kangar, Perlis, Malaysia

Mallinckrodt Développement France S.A.S.

10 Allee Pelletier Doisy, 54601 Villers-Lès-Nancy, France

Mallinckrodt DAR S.R.L.

Via G. Bove, 2/4/6/8, Mirandola MO 41037, Italy

Mallinckrodt Medical

Cornamaddy Athlone, Co. Westmeath, Ireland

SOMANETICS CORP.

1653 East Maple Road, Troy, MI 48083-4208, USA

Aspect Medical Systems

1 Upland Road, Norwood MA 02062, USA

POLYSUTURE

Av. Gabriel Ramos da Silva, 1245 Pq. Industrial II, São Sebestio. Paraíso, Minas Gerais, Cep: 37950-000 Brasil

D00928740 Revision AA Page 1 of 4



Declaration of Conformity

Legal Manufacturer: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

EC Authorized Representative Medtronic Ireland

Parkmore Business Park West

Galway Ireland

Design Facility: Medtronic Ireland

Parkmore Business Park West

Galway Ireland

Manufacturing Facility: Medtronic Ireland

Parkmore Business Park West

Galway, Ireland

Product Family/ies: Coronary and Peripheral Guide Catheters

Products: Telescope Guide Extension Catheter

See attached list of model numbers

Classification: Class III based on Annex IX, Rule 6 of Directive

93/42/EEC (MDD)

Notified Body British Standards Institution (BSI) 2797

EC Quality Certificate CE84868 first issued 24 August 2004

EC Design Certificate CE701802

Declaration of Conformity Form Medtronic D00928740 Page 2 of 4 Revision AA

I, the undersigned, hereby declare that the Medical Device(s) specified above meet the provisions of Article 120 of Medical Device Regulation (MDR) 2017/745, which replaced Directive 93/42/EEC (MDD), including amendments issued, on the date of application (May 26th, 2021).

This declaration is supported by Article 120 MDD 2017/745 and the MDD, Annex II.3 Approval as well as the EC Design Examination Certificate listed above (if applicable).

Signed on behalf of the manufacturer exclusively responsible for the Declaration of Conformity:

Medtronic Ireland

Date: 9th Feb, 2023

Elaine O'Connor
Senior Regulatory Affairs Manager Signature Elaine O'Connoc Place:

Name:

Title

Declaration of Conformity D00928740 Revision AA Page 2 of 4 Form Medtronic

I, the undersigned, hereby declare that the Medical Device(s) specified above meet the provisions of Article 120 of Medical Device Regulation (MDR) 2017/745, which replaced Directive 93/42/EEC (MDD), including amendments issued, on the date of application (May 26th, 2021).

This declaration is supported by Article 120 MDD 2017/745 and the MDD, Annex II.3 Approval as well as the EC Design Examination Certificate listed above (if applicable).

Signed on behalf of the manufacturer exclusively responsible for the Declaration of Conformity:

Place:	Medtronic Ireland	_ Date:	
Name:	Elaine O'Connor		
Title	Senior Regulatory Affairs Manager	Signature	

Declaration of Conformity D00928740 Revision AA Page 3 of 4 Form Medtronic

Products: Telescope Guide Extension Catheter

Model Name	Model No.
Telescope	TELE6F
Telescope	TELE7F

Declaration of Conformity





Revision History

Revision	Date	Description of Change
1A	04 April 2019	First Issue
AA	08 February 2023	Addition of the Optimised PCA sterilisation cycle





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

No. CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

In respect of:

The design, development and manufacture of sterile Endoluminal Stent Grafts, sterile Securement Devices and Delivery Systems for Endovascular Indications, sterile Vascular Introducer Sheaths, sterile Stent Graft Balloon Catheters, sterile Coronary Stents and Delivery Systems, Sterile Intravascular Catheters and sterile/non-sterile Catheter Systems for Renal Denervation.

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):

Gary E Slack, Senior Vice President - Medical Devices

Jany C Stade

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 1 of 6

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.





Supplementary Information to CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Number	Device Name	Intended purpose per IFU	
Class III pro	ducts under the scope of CE 84868		
N/A	Attain Clarity Venogram Balloon Catheter	See CE 593123	
N/A	Driver Sprint Rapid Exchange Coronary Stent System	See CE 545439	
N/A	Endeavor Resolute Zotarolimus-Eluting Coronary Stent System Resolute Integrity Zotarolimus-Eluting Coronary Stent System	See CE 514336	
N/A	Endeavor Sprint Zotarolimus-Eluting RX Coronary Stent System See CE 86406		
N/A	Endurant™ Stent Graft System Endurant™ II Stent Graft System Endurant™ IIs Stent Graft System		
N/A	Euphora Rapid Exchange Balloon Dilatation Catheter See CE 622066		
N/A	Heli-FX™ EndoAnchor™ Systems See CE 669930		
N/A	IN.PACT Admiral (Paclitaxel-coated PTA Balloon Catheter) See CE 570280		

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 2 of 6





Supplementary Information to CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Number	Device Name	Intended purpose per IFU	
Class III prod	lucts under the scope of CE 84868		
N/A	IN.PACT Falcon (Paclitaxel-eluting PTCA Balloon Catheter)	See CE 570282	
N/A	IN.PACT Pacific (Paclitaxel-eluting PTA Balloon Catheter)	See CE 570281	
N/A	Integrity Rapid Exchange Coronary Stent System	See CE 91271	
N/A	Micra™ Introducer Sheath with Hydrophilic Coating	See CE 599898	
N/A	NC Euphora Rapid Exchange Balloon Dilatation Catheter	See CE 612356	
N/A	NC Solarice Rapid Exchange Balloon Dilatation Catheter See CE 630635		
N/A	NC Sprinter Rapid Exchange Balloon Dilatation Catheter See CE 506473		
N/A	Reliant Stent Graft Balloon Catheter	See CE 635936	
N/A	Resolute Onyx Zotarolimus-Eluting Coronary Stent System See CE 618060		
N/A	Sentrant Introducer Sheath with Hydrophilic Coating See CE 595294		
N/A	Solarice Rapid Exchange Balloon Dilatation Catheter See CE 630580		
N/A	Sprinter Legend OTW Balloon Dilatation Catheter See CE 547584		

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 3 of 6





Supplementary Information to CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Number	Device Name	Intended purpose per IFU
Class III prod	ucts under the scope of CE 84868	
N/A	Sprinter Legend RX Balloon Dilatation Catheter	See CE 525652
N/A	Sprinter Over-the-Wire Balloon Dilatation Catheter See CE 92065	
N/A	Telescope Guide Extension Catheter See CE 701802	
N/A	Valiant Navion™ Thoracic Stent Graft System See CE 702496	
N/A	Valiant Thoracic Stent Graft with the Captivia Delivery System See CE 554030	
N/A	Prevail Paclitaxel-coated PTCA Balloon Catheter See CE 718244	

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 4 of 6





Supplementary Information to CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Class IIb products under the scope of CE 84868		
GMDN #	Device or Generic Device Group	Intended Purpose per IFU
58893 (Catheter) 35156 (Generator)	Symplicity Spyral [™] Multi-Electrode Renal Denervation Catheter & Symplicity G3 [™] Renal Denervation RF Generator	The Symplicity G3™ Renal Denervation RF Generator when used with the Symplicity Spyral™ Multi-Electrode Renal Denervation Catheter is intended to deliver low-level radio frequency (RF) energy through the wall of the renal artery to denervate the human kidney.

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 5 of 6





Supplementary Information to CE 84868

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Class IIb products under the scope of CE 84868			
GMDN #	Device or Generic Device Group	Intended Purpose per IFU	
46777	Talent Endoluminal Occluder System	The Talent Endoluminal Occluder System is intended for endoluminal occlusion of the contralateral iliac artery in cases where an abdominal aortic aneurysm is treated with an aorto-uni-iliac stent graft and subsequent femoral- to-femoral bypass procedure	

First Issued: **2004-08-24** Date: **2020-12-23** Expiry Date: **2024-05-26**

...making excellence a habit."

Page 6 of 6





Certificate No: **CE 84868**Date: **2020-12-23**

Issued To: **Medtronic, Inc.**

710 Medtronic Parkway Minneapolis, MN 55432

USA

Date	Reference Number	Action
24 August 2004		First Issued.
15 November 2004		Transfer of the following certificates from NSAI:-
		Q252.322, Q252.407, Q252.426, Q252.427, Q252.428, Q252.467, Q252.480, Q252.587, and Q252.611
		D252.587 and D252.407, plus incorporation of Medtronic Vascular Ireland as a subcontract manufacturer.
02 December 2004		Carotid and Coronary Stents and Delivery Systems added to the scope (transfer) Medtronic Mexico (manufacture), and Titan Scan Systems, Nutec Corporation, Sterigenics (Queensbury), Steris Corporation-Isomedix Services (Sandy), Rocialle in Health (Mid Glamorgan UK), and EBIS Iotron added as sub-contract sterilizers.
21 December 2004		PTCA Balloon Dilatation Catheters added to the range of products manufactured (transferred from another Notified Body) and Isotron Ireland Ltd added as sub-contract sterilization site.
19 August 2005		Sterilization sub-contractor name change from Titan Scan Systems to Beam One.
03 April 2006		Addition of Sterigenics UK Ltd, as sterilization sub-contractor.

...making excellence a habit."

Page 1 of 6





Certificate No: **CE 84868**

Date: 2020-12-23
Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Date	Reference Number	Action	
07 August 2006		Addition of AD)MEDES Schuessler GmbH as a sub-contractor for manufacture.	
11 January 2008	7149866	Subcontractor name change from EBIS Isotron, Harwell to Isotron Harwell. Addition of Isotron plc, Daventry as a subcontractor for E beam sterilization.	
03 October 2008	7279045	Addition of Medtronic Mexico EG, Empalme as a subcontractor for manufacture.	
14 April 2009	7341499	Correction of the legal name of the Medtronic Mexico facility and postcode for the Isotron PLC, Daventry facility. Addition of the activity of EU Representative for Medtronic Ireland.	
13 August 2009	7432878	Addition of the activity of EU Representative for Medtronic Ireland. Certificate renewal. Addition of Accellant Inc as a manufacturing subcontractor, amendment to company name for Isotron PLC, Daventry, and Steris Corporation, Sandy, Utah. Change to address for the subcontractor, Nutek Corporation. Addition of E Beam Sterilization for Isotron Ireland. Rewording of scope for clarification purposes only.	

...making excellence a habit.™

Page 2 of 6

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.





Certificate No: **CE 84868**

Date: 2020-12-23

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Date	Reference Number	Action
29 July 2010	7546410	Added C.R. Bard, Inc. to the list of significant subcontractors for manufacturing. Extended the scope to include guidewires.
12 October 2011	7730209	Extension to scope to include Catheter Systems for Renal Denervation. Removal of Carotid Stents and Delivery Systems from the scope. Minor amendments to Isotron Daventry and Isotron Tullamore's addresses.
26 January 2012	7792125	Amendment to significant subcontractors to reflect Isotron's name change to Synergy Health and removal of Isotron Harwell.
25 May 2012	7842435	Amendment to the address format and zip code for the significant subcontractor Medtronic Mexico (Tijuana).
19 December 2012	7915649	Addition of Medtronic B.V. The Netherlands for EU Representative Activities.
22 January 2013	7945194	Extension to scope to include Superficial Femoral Artery (SFA) and Proximal Popliteal Artery (PPA) Stents and Delivery Systems.
28 February 2013	7960715	Addition of Invatec Technology Center GmbH to the list of significant subcontractors for manufacturing activities.
28 March 2013	7943883	Extension to Scope to include Vascular Introducer Sheaths and the addition of Teleflex Medical for manufacturing activities.

...making excellence a habit.™

Page 3 of 6

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.





Certificate No: **CE 84868**Date: **2020-12-23**

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Date	Reference Number	Action	
16 December 2013	8082854	Addition of Plexus Manufacturing Sdn Bhd, Malaysia and Plexus Corp, UK to the list of significant subcontractors for manufacturing activities.	
13 July 2014	8154862	Certificate Renewal. Various updates and changes to the list of significant subcontractors. Correction of the reference number for the reissue dated 19 th December 2012 on the certificate history page.	
31 July 2015	8350802	Addition of SSP SiMATrix Inc. as balloon supplier for the Attain Clarity.	
01 July 2016	8545838	C. R. Bard, Inc., Medtronic Ardian LLC, Nutek Corporation, Sterigenics NY and Apical Instruments Inc. were removed from the list of significant subcontractors.	
09 October 2017	8696759	Certificate scope updated to add the design, development and manufacture of securement devices for endovascular indications.	
01 May 2018	8895951	Specify devices covered in this certificate are sterile/non-sterile. Move 'sterile Vascular Introducer Sheaths' up in the scope after securement devices. Remove 'Renal Stents and Delivery Systems' and 'guidewires for diagnostic or interventional procedures' from scope. Correction to certificate history entry #2 from '2014' to '2004'.	
06 March 2019	8786554	Traceable to NB 0086.	

...making excellence a habit."

Page 4 of 6

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.





Certificate No: **CE 84868**Date: **2020-12-23**

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Date	Reference Number	Action
22 August 2019	9736517	Certificate Renewal.
		Added product table per MDP4500 Appendix A.
		Clarified addresses of subcontractors to exactly align with their ISO certificate name and address.
		Remove "sterile Iliac Stents and Delivery Systems, sterile Superficial Femoral Artery (SFA) and Proximal Popliteal Artery (PPA) Stents and Delivery Systems" from scope as the Complete SE product (iliac and vascular indications) is no longer manufactured nor in the distribution chain.
		Remove Assurant Cobalt product (iliac product scope) it is no longer manufactured and the last product builds expired in April 2019.
		Remove subcontractors – Admedes Schuessler GmbH, Germany, Flextronics Medical, Austria, Sterigenics, Corona, CA, Synergy Health, Ireland related to removed products above.
		Add subcontractors - Phoenix DeVentures, CA, Sterigenics, Los Angeles, CA, SurModics, MN and Medtronic, Santa Ana, CA related to new Class IIa product Confida Expandable Sheath.
09 October 2020	3295912	Remove Class IIa Confida Expandable Sheath from product table. Add Class III Prevail Paclitaxel-coated PTCA Balloon Catheter to product table. Add crucial supplier Indena S.p.A. Remove subcontractors Phoenix DeVentures, Inc., Sterigenics US, LLC, Medtronic Corevalve LLC and SurModics, Inc. related to the Confida Expandable Sheath.

...making excellence a habit."

Page 5 of 6

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.





Certificate No: **CE 84868**

08 October 2021

01 February 2023

2020-12-23 Date:

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

Date	Reference Number	Action	
23 December 2020	3329282	Addition of significant subcontractor Avantti Clear MBP, S.A.P.I C.V., Tijuana, Mexico for the service of Radiation (E-Beam Sterilization) for the NC Solarice, NC Sprinter and NC Euphora devices.	
Non-significant cl of MDR Article 12		ed after the 26 th May 2021 as per the Transitional Provisions	
04 August 2021	3497705	Remove "Endeavor Sprint Zotarolimus-Eluting RX Coronary Stent System – See CE 86406" and "NC Sprinter Rapid Exchange Balloon Dilatation Catheter – See CE 506473" from the device table.	
08 October 2021	2515102	Addition of Medtronic Vascular (Danvers, MA) as a 'Manufacture'	

Remove all subcontractor pages.

...making excellence a habit."

Page 6 of 6

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.

3515102

3834700

supplier.



Inspiring trust for a more resilient world.

01 February 2023 Medtronic, Inc. 710 Medtronic Parkway Minneapolis, MN 55432 USA

To whom it may concern,

The transitional provisions specified in MDR Article 120(3) prohibit Notified Bodies from issuing new certificates or amending, modifying, supplementing any existing MDD/AIMDD certificates from 26th May 2021. This letter is to confirm that BSI has reviewed and approved the change(s) detailed in the table below. These changes do not represent a significant change in design or intended purpose under MDR Article 120(3) and as per the guidance provided in MDCG 2020-3. The related MDD certificate specified below remains valid until the expiry date specified on the certificate.

Certificate	Directive and Annex	Reference Number	Changes approved
CE 84868	93/42/EEC Annex II excluding Section 4	3834700	Addition of a critical subcontractor and administrative update for removal of all subcontractor pages

Should you have any queries concerning your certification, or if we can be of further assistance to you, please contact your BSI Scheme Manager.

Yours sincerely,

Graeme Tunbridge

Senior Vice President, Medical Devices







EC Design-Examination Certificate

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

CE 701802

Issued To: Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

In respect of:

Telescope Guide Extension Catheter

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 2797):

Gary E Slack, Senior Vice President Medical Devices

First Issued: **2019-04-04** Date: **2019-04-04** Expiry Date: **2024-04-03**

...making excellence a habit."

Page 1 of 3

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

Gay C Stade





EC Design-Examination Certificate

Supplementary Information to CE 701802

Issued To:

Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Product Listing

Telescope Guide Extension Catheter

Catalogue Number	Device Name	French Size	Intended Purpose per IFU	Classification
TELE6F		6 Fr	Telescope Guide Extension Catheter is intended to be used in conjunction with guide catheters to access discrete regions of the coronary and/or peripheral vasculature, and to facilitate placement of interventional devices.	Class III
TELE7F	Telescope Guide Extension Catheter	7 Fr		Class III

First Issued: **2019-04-04** Date: **2019-04-04** Expiry Date: **2024-04-03**

...making excellence a habit."

Page 2 of 3

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.





EC Design-Examination Certificate

Supplementary Information to CE 701802

Issued To:

Medtronic, Inc.

710 Medtronic Parkway Minneapolis, MN 55432

USA

Certificate History

Date	Reference Number		Action	
Current	9666412	First Issue.		100

First Issued: **2019-04-04** Date: **2019-04-04** Expiry Date: **2024-04-03**

...making excellence a habit."

Page 3 of 3

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 701802 - Non-significant changes approved after the 26th May 2021 as per the Transitional Provisions of MDR Article 120.3

Issued to: **Medtronic, Inc.**

710 Medtronic Parkway Minneapolis MN 55432

USA

Date: 18 August 2021

Changes Approved:

Date	Reference Number	Action
18 August 2021	3443630	Addition of optimized PCA EO sterilization cycle.



Inspiring trust for a more resilient world.

18 August 2021

Medtronic, Inc. 710 Medtronic Parkway Minneapolis, MN 55432 USA

To whom it may concern,

The transitional provisions specified in MDR Article 120(3) prohibit Notified Bodies from issuing new certificates or amending, modifying, supplementing any existing MDD/AIMDD certificates from 26th May 2021.

This letter is to confirm that BSI has reviewed and approved the change(s) detailed in the table below. These changes do not represent a significant change in design or intended purpose under MDR Article 120(3) and as per the guidance provided in MDCG 2020-3. The related MDD certificate specified below remains valid until the expiry date specified on the certificate.

Certificate	Directive and Annex	Reference Number	Changes approved
CE 701802	93/42/EEC Annex II Section 4	3443630	Addition of optimized PCA EtO Sterilization cycle to reduce EO concentration.

Should you have any queries concerning your certification, or if we can be of further assistance to you, please contact your BSI Scheme Manager.

Yours sincerely,

Gary Slack

Senior Vice President, Medical Devices

jany C Stade



