



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

No. Issued To: CE 540595 Teleflex Medical IDA Business and Technology Park Dublin Road Athlone 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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Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

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

Information and Contact: BSI, Kitemark Court, Davy Avenue, Knowlhill, Milton Keynes MK5 8PP. Tel: + 44 845 080 9000 BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK. A member of BSI Group of Companies.





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:

CE 540595

Certificate No: Date: Issued To:

28 August 2015 Teleflex Medical IDA Business and Technology Park Dublin Road Athlone Co. Westmeath Ireland

Subcontractor:

Arrow International CR, a.s. Jamska 2359/47 59101 Zdar nad Sazavou Czech Republic

Arrow International CR, a.s. Prazska 209 50004 Hradec Kralove Czech Republic

Arrow Medical Ltd Hatton Gardens Industrial Estate Kington HR5 3RB United Kingdom

CeMed GmbH Oberdorf 41 72419 Neufra Germany Service(s) supplied

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Page 1 of 5





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Subcontractor:

Service(s) supplied

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Chelle Medical Limited PO Box 221 Le Rocher Victoria Mahe Seychelles

Forefront (Xiamen) Medical Devices Co., Ltd No 26 & 28 Haijing Dong Lu Haicang Xiamen Export Processing Zone 361026, Xiamen, Fujian China

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

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Page 2 of 5





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Subcontractor:

Service(s) supplied

M.E.M., Inc. 8 Bishop Lane Madison Connecticut 06443 USA

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**

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Page 3 of 5





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Ireland

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Subcontractor:

SP Medical A/S Møllevej 1 4653 Karise Denmark

Süddeutsche Feinmechanik GmbH (SFM) Brückenstrasse 5 D-63607 Wächtersbach Germany

Teleflex Medical Sdn. Bhd. Lot PT2577, Jalan Perusahaan 4 34600 Kamunting Perak Malaysia

Teleflex Medical Asia Pte. Ltd. 6 Battery Road #07-02 049909 Singapore Service(s) supplied

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Page 4 of 5





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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

Tianjin Medis Medical Device Co. Ltd 10A Tianzhi Industrial Centre No 12 Hong Yuan Road Xiqing Economic Development Area 300385 Tianjin City China

Willy Rüsch GmbH Willy Rüsch-Strasse 4-10 D-71394 Kernen Germany **Crucial Supplier**

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Page 5 of 5





EC Certificate - Full Quality Assurance System Certificate History

Certificate No: Date: Issued To: CE 540595 28 August 2015 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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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.

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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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