

EC Certificate - Full Quality Assurance System

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

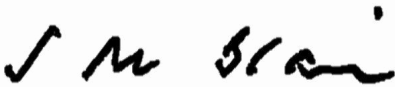
No. CE 584854
Issued To: **DySIS Medical Limited**
Gyleview House
3 Redheughs Rigg
Edinburgh
EH12 9DQ
United Kingdom

In respect of:

The design and manufacture of digital colposcopy device for quantified mapping of the aceto-whitening effect on the vagina, cervix and external genitalia.
Those aspects of Annex II relating to securing and maintaining sterility in the design and manufacture of the sterile single use specula and of the sterile single use disposable treatment pipe.

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: **2012-07-02**

Date: **2017-06-16**

Expiry Date: **2022-07-01**

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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.

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List of Significant Subcontractors

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

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Subcontractor:	Service(s) supplied
Andersen Caledonia Limited Caledonian House Phoenix Crescent Strathclyde Business Park North Lanarkshire Scotland ML4 3NJ United Kingdom	Sterilization
Eurolaz Technologies Ltd The Maltings Industrial Estate Hall Road, Southminister CM0 7EQ United Kingdom	Control of Sterilization Sterile Manufacture
Fearsom UK Ltd (t/a Fearsome) The Whisky Bond 2 Dawson Road Glasgow G4 9SS United Kingdom	Design

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Edinburgh
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Subcontractor:	Service(s) supplied
ITL Viking House Elingham Way Ashford, Kent TN23 6NF United Kingdom	Design Manufacture
RB Medical Engineering Ltd Alton Road Industrial Estate Ross-on-Wye Herefordshire HR9 5NS United Kingdom	Manufacture
Sanmina-SCI AB Svedjevagen 12 Sjalevad 894 35 Sweden	Design Manufacture

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Date	Reference Number	Action
02 July 2012	7806173	First issue
04 August 2014	8134668	Change of scope adding "including sterile and non-sterile specula". Addition of subcontractor Europlaz Technologies Ltd for "Sterile Manufacture".
30 April 2015	8333446	Removal of subcontractor DySIS Medical Hellas.
03 October 2016	8575902	Addition of subcontractor Fearsom UK Ltd (t/a Fearsome) for Design. Change of address of the legal manufacturer from Alba Innovation Centre, Livingston, EH54 7GA to Westpoint, 4 Redheughs Rigg, Edinburgh EH12 9DQ. Change of scope from "The design and manufacture of digital colposcopy devices for quantified mapping of the aceto-whitening effect on the vagina, cervix and external genitalia, including sterile and non-sterile specula. " to reflect the reclassification of the specula.

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Current	8747443	<p>Renewal.</p> <p>Change of address from DYSIS Medical Limited, Westpoint, 4 Redheughs Rigg, Edinburgh, EH12 9DQ to DYSIS Medical Limited, Gyleview House, 3 Redheughs Rigg, Edinburgh, EH12 9DQ.</p> <p>Addition of service supplied "Control of Sterilization" for subcontractor Europlaz Technologies Ltd.</p> <p>Addition of subcontractor Andersen Caledonia Ltd for "Sterilization".</p> <p>Addition of subcontractor Sanmina-SCI AB for "Design and Manufacture".</p> <p>Change of scope to include DYSIS Disposable Treatment Pipe.</p>
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