

Quality System Approval Certificate

Medical Devices Directive 93/42/EEC

The National Standards Authority of Ireland as a duly designated Notified Body, (identification number **0050**), for the purposes of the European Communities (Medical Devices) Regulations (S.I. No. 252 of 1994)

APPROVES THE QUALITY SYSTEM APPLIED BY

Lumitex Medical Devices, Inc.

8443 Dow Circle Strongsville Ohio 44136 USA

to the Product Family

Surgical instrument fibreoptic light (LightMat)

GMDN Code: 48036

on the basis of examination under the requirements of Directive 93/42/EEC on Medical Devices, Annex II, excluding (4)

The use of the NSAI Notified Body identification number **0050** in conjunction with CE Marking of Conformance for this product family is hereby authorised.

Registration Number:

Original Approval:

Last Amended on:

Remains valid until:

Signed:

Approved by: Geraldine Larkin Chief Executive Officer, NSAI 252.914

14 November 2013

19 December 2017

09 April 2018

Approved by: Susan Murphy European Medical Device Operations Manager

This certificate remains valid on condition that the Approved Quality System is maintained in an adequate and efficacious manner. Details of the operational locations included within the scope of this approval can be obtained from NSAI

In the case of a Class III device, this certificate must be supported by a valid design examination certificate

National Standards Authority of Ireland, 1 Swift Square, Northwood, Santry, Dublin 9, Ireland.