



# NSAI

## 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.

**8300 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.*

|                             |                          |
|-----------------------------|--------------------------|
| <b>Registration Number:</b> | <b>252.914</b>           |
| <b>Original Approval:</b>   | <b>14 November 2013</b>  |
| <b>Last Amended on:</b>     | <b>03 March 2021</b>     |
| <b>Remains valid until:</b> | <b>24 September 2023</b> |

**Signed:**

Approved by:  
Dr. Caroline Dore Geraghty  
Director, Medical Devices

Approved by:  
Dr. Elaine Darcy  
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.**