

Certificate of Registration of Quality Management System to I.S. EN ISO 13485:2016

The National Standards Authority of Ireland certifies that: Lumitex Medical Devices, Inc. 8300 Dow Circle Suite 400 Strongsville, OH 44136 USA

has been assessed and deemed to comply with the requirements of the above standard in respect of the scope of operations given below:

The Design and Development, Manufacture and Distribution of Sterile and Non-Sterile Medical Illumination Devices and Nonsterile Phototherapy Devices.

Additional sites covered under this multi-site certification are listed on the Annex (File No. MD19.4787/A)

Approved by: Kevin Mullaney Director of Certification

Registration Number: MD19.4787/A Certification Granted: March 26, 2013 Effective Date: June 26, 2023 Expiry Date: June 25, 2026



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Annex to Certificate Number: MD19.4787/A

Scope of Registration:

The Design and Development, Manufacture and Distribution of Sterile and Non-Sterile Medical Illumination Devices and Nonsterile Phototherapy Devices.

Activity

Headquarters, Quality Assurance, Regulatory Affairs, Production, and Design and Development

Quality Assurance, Regulatory Affairs, Production, Design and Development

Location

Lumitex Medical Devices, Inc. 8300 Dow Circle Suite 400 Strongsville, OH 44136 USA File No.: MD19.4787/A

Lumitex Medical Devices, Inc. 8443 Dow Circle Strongsville, OH 44136 USA File No.: MD19.4787

Verified by: Director of Certification