

Certificate

acc. to **ISO 13485:2016**

Medical Devices - Quality Management Systems - Requirements for Regulatory Purposes

Certificate Registration No.: 23-1603-Q

TUV USA, Inc. hereby certifies that the quality management system of the company mentioned below is in conformance with **ISO 13485:2016** for Medical Devices-Quality Management Systems-Requirements for Regulatory Purposes.



Weinert Fiber Optics, Inc.
209 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

Additional sites covered by QM System: [See Annex 1](#)

Scope:

Design, Development, Production, and Distribution of Sterile and Non-Sterile Fiber Optic Devices, Components, and Assemblies for Medical Applications

The validity of this certification document can be obtained by contacting the TUV USA, Inc. Office.

TUV USA, Inc. (a Member of the TÜV NORD Group)

215 Main Street, Suite 1, Salem, NH 03079, USA

Tel: 001-603-870-8023, Fax: 001-603-870-8026, Email: medical-usa@tuv-nord.com



Audit Report Reference No.: 23-4050 CA

Initial Certification Date: 2023-08-15

Current Cycle Start Date: 2023-08-15

Effective Date:
2023-08-15 / ed. 1

Valid Until:
2026-08-14

Bradley Chen
Vice President – Medical, Americas
Medical Products Division
TUV USA, Inc.

Certificate Registration No. : 23-1603-Q / ed. 1
Company Name: Weinert Fiber Optics, Inc.
Central Office Address: 209 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

Additional Sites covered by the QM System:

Location

Scope of Certification

Headquarters

Weinert Fiber Optics, Inc.
209 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

HQ, Top Management, Manufacturing,
Purchasing, and HR

Site 01

Weinert Fiber Optics, Inc.
203 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

Manufacturing, Design, Sales, and
Document Control

Site 02

Weinert Fiber Optics, Inc.
215 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

Manufacturing and Sterilization

Site 03

Weinert Fiber Optics, Inc.
219 Bulifants Blvd.
Williamsburg, Virginia 23188, USA

Manufacturing (Non-Medical)

---End of list---

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A handwritten signature in blue ink, reading "Bradley Chen".

Bradley Chen
Vice President – Medical, Americas
Medical Products Division
TÜV USA, Inc.