

EC Certificate

Full Quality Assurance System according to Medical Devices Directive 93/42/EEC Annex-II Section 3

Certificate Number: 1984-MDD-16-372

We hereby declare that an examination of the under mentioned full quality assurance system has been carried out following the requirements of the national legislation to which the undersigned is subjected, transposing annex II (with the exemption of section 4) of the Directive 93/42/EEC on medical devices. We certify that the full quality assurance system conforms with the relevant provisions of the aforementioned directive.

Organization:

CeramOptec GmbH

Siemensstrasse 44, 53121 Bonn, Germany
Facility: Brühler Strasse 30 53119 Bonn, Germany

Products: The products defined at the enclosure which is the part of this certificate and contains two pages.

The certificate is valid till expiration date, subject to successful completion of periodical surveillance audits. Please contact Kiwa for details.

Report Number: M.4508.05
Date of first issue: 14 March 2016
Date of last issue: 04 September 2020
Revision Number: 03
Expiry Date: 12 March 2024



Muhteşem Gökhan Yücel
Head of Notified Body

04 September 2020, Istanbul, Turkey

Enclosure of the Certificate:**Full Quality Assurance System according to****Medical Devices Directive 93/42/EEC Annex II Section 3****Certificate Number: 1984-MDD-16-372, Revision Number: 03**

Concerned medical devices;

Product: Diode Lasers

- Types:**
- Type Ceralas E
 - Type Ceralas HPD
 - Type Leonardo
 - Type Leonardo HPD
 - Type Leonardo Mini
 - Type Leonardo Bonsai

Product: Probes for Lasers

- Types:**
- Type Bare Fiber, single-use, sterile
 - Type Bare Fiber, reusable, sterile
 - Type Endoprobe, single-use, sterile
 - Type Gas Liquid Cooled, single-use, sterile
 - Type Side Fiber, single-use, sterile
 - Type PLDD Bare Fiber, single-use, sterile
 - Type Cylindrical diffuser, single-use, sterile
 - Type ELVeS Fiber, single-use, sterile
 - Type Twister, single-use, sterile
 - Type ELVeS Radial, single-use, sterile
 - Type Bare fiber for Ho:YAG Laser, single-use, sterile
 - Type Bare fiber for Ho:YAG Laser, reusable, sterile

A handwritten signature in black ink, appearing to read "Muhteşem Gökhan Yücel".

Muhteşem Gökhan Yücel
Head of Notified Body

04 September 2020, Istanbul, Turkey

Enclosure of the Certificate:

Full Quality Assurance System according to

Medical Devices Directive 93/42/EEC Annex II Section 3

Certificate Number: 1984-MDD-16-372, Revision Number: 03

Concerned medical devices;

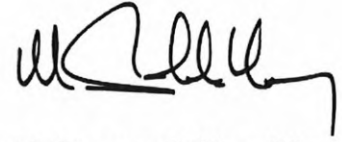
Product: Handpieces

Type: Type Derma Handpiece; reusable

Product: Introducer for Probes

Type: Type ELVeS Plus Catheter, sterile

Kiwa Belgelendirme Hizmetleri A.Ş. is Notified Body under Council Directive 93/42/EEC concerning medical devices with identification number: 1984



Muhteşem Gökhan Yücel
Head of Notified Body

04 September 2020, Istanbul, Turkey

CERTIFICATE