



EU Quality Management System Certificate
Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III
Certificate No. MDR-001

Issued to: Meridian Medical d.o.o.
Plemljeva ulica 8, 1210 Ljubljana-Šentvid
Slovenija

SRN of the manufacturer: SI-MF-000006685

EU authorised representative: Not applicable

SRN of EU authorised
representative: Not applicable

SIQ has audited the quality management system in accordance with MDR Annex IX and found that the above-mentioned Manufacturer's quality management system meets the requirements of the Regulation (EU) 2017/745 concerning medical devices Annex IX. Devices covered by the Manufacturer's quality management system are listed on the page(s) below.
This certificate is based on

Audit report No.:

OSV 01133/2022, 2022-09-30

OSV 01548/2022, 2022-12-30

OSV 00316A/2023, 2023-03-23

See also decision of NB's commission for medical devices.

This certificate remains valid as long as the Manufacturer's quality management system is subject to periodical surveillance as referred to in Regulation (EU) 2017/745 concerning medical devices Annex IX and continues to meet the above requirements.

Reference to any previous certificate: /

Certification date: 2023-03-23

Issue: 01/2023-03-23

Valid until: 2028-03-22

Managing Director of SIQ

Gregor Schoss





EU Quality Management System Certificate
Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III
Certificate No. MDR-001

Device: Neodymium surgical laser

EMDN: Z12011017

Intended purpose: MR Q SLT is an ophthalmic medical device that provides the ability to treat patients in a sitting position using Nd:YAG laser source with 1064 nm nanosecond pulsed output (YAG mode) or with 532 nm (KTP frequency-doubled) nanosecond pulsed output (SLT mode) regarding to selected treatment.

Classification: IIb

Specific conditions for or /
provisions or limitations to the
validity of certificate:



Z vami več kot
With you for over
50 let
years

SIQ Ljubljana

Mašera-Spasičeva ulica 10
SI-1000 Ljubljana
Slovenija

T +386 1 4778 100
F +386 1 4778 444
info@siq.si, www.siq.si

Mr.
Kristijan Lamot
Meridian Medical d.o.o.
Plemljeva ulica 8

1210 LJUBLJANA-ŠENTVID

vaš znak / your ref.: OSV/5361
naš znak / our ref.: 2023-03-23
datum / date:

Dear Mr Lamot!

We would like to inform you that the Notified Body Commission for medical devices adopted the following decision at its last meeting:

On the basis of certification audit report (OSV 01133/2022 dated 2022-09-30), technical documentation audit report and technical documentation post audit report (OSV 01548/2022 dated 2022-12-30) and technical documentation post-audit report and post audit report (OSV 00316A/2023 dated 2023-03-23), a decision on granting the EU certificate was adopted:

EU certificate No.: **MDR-001**

Issued to:	Meridian Medical d.o.o. Plemljeva ulica 8, 1210 Ljubljana-Šentvid Slovenija
SRN of manufacturer:	SI-MF-000006685
Authorised representative:	Not applicable
SRN of authorised representative:	Not applicable

Legislative act/requirements: Regulation (EU) 2017/745
Annex: IX

Device:	Neodymium surgical laser
EMDN:	Z12011017
Intended purpose:	MR Q SLT is an ophthalmic medical device that provides the ability to treat patients in a sitting position using Nd:YAG laser source with 1064 nm nanosecond pulsed output (YAG mode) or with 532 nm (KTP frequency-doubled) nanosecond pulsed output (SLT mode) regarding to selected treatment.
Classification:	IIB

Certification date: 2023-03-23
Issue: 01/2023-03-23
Valid until: 2028-03-22

Specific conditions or provisions or limitations and validity of certificate: /

Identification Number of the Notified Body (1304) can be used in conjunction with EU certificate. Details of affixing CE marking are given in Article 20 of Regulation (EU) 2017/745 on medical devices.

The EU certificate (A4 paper format) is granted in English. EU certificate is enclosed in attachment. Also form for possible additional order of certificate is enclosed.

Detailed list of product names, models and types:

EU certificate No.: **MDR-001**
Issued to: Meridian Medical d.o.o.
Plemljeva ulica 8, 1210 Ljubljana-Šentvid
Slovenija
SRN of manufacturer: SI-MF-000006685
Place of production: Meridian Medical d.o.o.
Plemljeva ulica 8, 1210 Ljubljana-Šentvid
Slovenija
Authorised representative: Not applicable
SRN of authorised representative: Not applicable
Device: Neodymium surgical laser
EMDN: Z12011017
Intended purpose: MR Q SLT is an ophthalmic medical device that provides the ability to treat patients in a sitting position using Nd:YAG laser source with 1064 nm nanosecond pulsed output (YAG mode) or with 532 nm (KTP frequency-doubled) nanosecond pulsed output (SLT mode) regarding to selected treatment.
Basic UDI-DI: 383007498MTLSXX1XG
MDR code: MDA 0302
Classification: IIb
Product name: MR Q SLT
Model/Type and/or Reference/ Catalogue number: MR Q SLT s, MR Q SLT c
Legislative act/requirements: Regulation (EU) 2017/745
Date of issuing the list: 2023-03-23
Valid until: 2028-03-22

Detailed list of relevant common specifications and standards

EU certificate No.: **MDR-001**
Issued to: Meridian Medical d.o.o.
Plemljeva ulica 8, 1210 Ljubljana-Šentvid
Slovenija

SRN of manufacturer: SI-MF-000006685

Authorised representative: Not applicable

SRN of authorised representative: Not applicable

List of relevant common specifications and standards: EN 60601-1:2006/A2:2021 (IEC 60601-1:2005/A2:2020)
EN 60601-1-2:2015/A1:2021 (IEC 60601-1-2:2014/A1:2020)
EN 60601-1-6:2010+A1:2015+A2:2021 (IEC 60601-1-6:2010+AMD1:2013+AMD2:2020)
EN 60601-1-9:2008+A1+A2:2020 (IEC 60601-1-9:2007/A2:2020 (Rev 1))
EN IEC 60601-2-22:2020 (IEC 60601-2-22:2019)
EN 60825-1:2014/A11:2021 (IEC 60825-1:2014)
EN 62304:2006/A1:2015 (IEC 62304:2006/A1:2015)
EN ISO 14971:2019/A11:2021
EN ISO 13485:2016/A11:2021
EN 62366-1:2015/A1:2020 (IEC 62366-1:2015/A1:2020)
EN ISO 15004-2:2007
EN ISO 10993-1:2020 (ISO 10993-1:2018)
EN ISO 15223-1:2021
EN 207:2017

Device: Neodymium surgical laser
EMDN: Z12011017
Basic UDI-DI: 383007498MTLSXX1XG

Legislative act/requirements: Regulation (EU) 2017/745
Date of issuing the list: 2023-03-23
Valid until: 2028-03-22

For all further information we are available on telephone number 00 386 1 4778 159 and email: mdr@siq.si

Management Systems Assessment
MDR Product Manager
Ana Pribaković Borštnik

