

Certificate

The certification body

MEDCERT Zertifizierungs- und Prüfungsgesellschaft für die Medizin GmbH
Pilatuspool 2 – 20355 Hamburg – Germany

herewith certifies that the company

TERUMO EUROPE N.V.
Interleuvenlaan 40
3001 Leuven
Belgium

with locations listed in the appendix

has introduced, applies and maintains a quality management system in the area of:

**Design and development, manufacture and final inspection of
Stent systems**

The conformity of this quality management system to the requirements of the below mentioned standard was verified by an audit:

EN ISO 13485:2016

This certification is subject to surveillance by MEDCERT.

Effective date: 2019-12-19
Expiry date: 2022-12-18

Report No.: 3179FS15F
Procedure No.: QS – 3479
Certificate No.: 3479GB445191217

Hamburg, 2019-12-17



MEDCERT Certification Body
(Dr. Andreas Schich)

The certificate is only valid when provided entirely with all of its pages.
To verify the validity of this certificate, contact info@medcert.de.

MEDCERT is a DAkkS accredited management systems
certification body

Appendix of certificate

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List of locations included in the scope of certificate**Manufacturing site**

Ashitaka Factory of Terumo Corporation
150, Maimaigi-cho, Fujinomiya City
Shizuoka Prefecture 418-0015
Japan

Designing site

Terumo Corporation, R&D Center
1500, Inokuchi, Nakai-machi, Ashigarakami-gun,
Kanagawa Prefecture 259-0151
Japan

– End of list –

This appendix is integral part of the above-referenced certificate.
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