EC CERTIFICATE

for the Quality Assurance System



according the Directive 93/42/EEC, Annex II excluding section (4)

As a Notified Body of the European Union, DEKRA Certification GmbH certifies, that the company

Joline GmbH & Co. KG

Neue Rottenburger Straße 50, 72379 Hechingen, Germany

applies a quality assurance system according to the Directive 93/42/EEC Annex II for the medical devices listed in the annex. The approval is based on the result of the re-certification audit report no. 50565-Z5-00, the decision dated 2018-10-04 and is only valid in connection with the successful performance of the annual surveillance audits.

This certificate is valid from 2018-11-30 to 2023-11-29

Registration No.: 50565-16-06



DEKRA Certification GmbH Stuttgart; 2018-10-04

Notified Body ID-number: 0124



Benannt durch/Designated by

Zentralstelle der Länder & für Gesundheitsschutz & für Gesundheitsschutz & Medizinprodukten

ZLG-BS-295.10.02

Annex to the EC Certificate No. 50565-16-06

Valid from 2018-11-30 to 2023-11-29

Revision status of the annex: 1 dated 2019-05-20

Devices/device categories included in the certificate:

Class II a:

MD 0102

- Dialysis Catheter ST
 - Kits
 - Catheter

MD 0106

- Kyphoplasty Systems ALLEVO
 - Kits
 - Individual Instruments
- Dialysis Accessories
 - Introducer Needle
 - Guide Wire
 - Dilator
 - Trocar
 - Connector LT

Class III:

MD 0203

- Dialysis Catheter PU-LT
 - Kits
 - Catheter
- Dialysis Catheter Silicone LT
 - Kits
 - Catheter

MD 0106

Biopsy Forceps KNIPSA

For the placing on the market of class III devices covered by this certificate an EC design-examination certificate according to directive 93/42/EEC annex II (4) is required.

Ruth Delbeck-Bayer

DEKRA Certification GmbH, Stuttgart, 2019-05-20

Notified Body ID-number: 0124

EC CERTIFICATE

for the Quality Assurance System



according the Directive 93/42/EEC, Annex V

As a Notified Body of the European Union, DEKRA Certification GmbH certifies, that the company

Joline GmbH & Co. KG

Neue Rottenburger Straße 50, 72379 Hechingen, Germany

applies a quality assurance system according to the Directive 93/42/EEC Annex V for the medical devices listed in the annex. The approval is based on the result of the re-certification audit report no. 50565-Z5-00, the decision dated 2018-10-04 and is only valid in connection with the successful performance of the annual surveillance audits.

This certificate is valid from 2018-11-30 to 2023-11-29

Registration No.: 50565-17-05



DEKRA Certification GmbH Stuttgart; 2018-10-04

Notified Body ID-number: 0124

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Benannt durch/Designated by

Zentralstelle der Länder & für Gesundheitsschutz & hei Arzneimitteln und Medizinprodukten

ZLG-BS-295.10.02

Annex to the EC Certificate No. 50565-17-05

Valid from 2018-11-30 to 2023-11-29

Revision status of the annex: 0 dated 2018-11-30

Devices/device categories included in the certificate:

Class I s:

For the products listed below, review of the Quality Assurance System refers exclusively to aspects of manufacture concerned with securing and maintaining sterile conditions.

MD 0101

- Miniclamp

MD 0106

Mixer



DEKRA Certification GmbH, Stuttgart, 2018-10-04

Notified Body ID-number: 0124