Certificate

Production Quality Assurance System Approval Annex V of the Directive on Medical Devices

ECM, Bismarckstr.106, 52066 Aachen, notified to EC under 0481 hereby declares that an examination of the under mentioned quality assurance system has been carried out following the requirements of annex V of the Directive 93/42/EEC.



This certificate is issued on behalf of:

Manufacturer

intra special catheters GmbH Oststrasse 2, 66780 Rehlingen-Siersburg, Germany

ECM certifies that the quality assurance system under which the products listed in annex I to this certificate are manufactured conforms with the requirements of annex V of the Directive 93/42/EEC on medical devices.

This Certificate is only valid for the products mentioned above. Special terms of validity are described in annex I to this certificate.

Any substantial changes of the quality assurance system or the listed products which might affect conformity to annex V of the Directive 93/42/EEC have to be notified to ECM and are subject to a separate assessment.

Report Number Registered under Valid until

218-15-2018 Z/15/03696E November 17th, 2020

Aachen, November 18th, 2015

Certification Body



Annex I to Certificate Z/15/03696E

Number of Pages: 1 of 1



This certificate is valid for the hereafter following devices:

Name of product category	Name of individual type	Nomenclature code
Single use devices	Fittings, Adapter	11-726
Single use devices	Drainage Systems, Pleural	10-817
Single use devices	Drains, Thoracic	11-308
Single use devices	Guide Wires	11-925
Single use devices	Catheters, Introducers	10-678
Single use devices	Catheters, Vascular, Blood Pressure	10-689
Single use devices	Catheters, Vascular, Embolectomy/Thrombectomy	10-714
Single use devices	Catheters, Cardiac, Pericardium Drainage	10-741
Single use devices	Catheters, Others	15-209
Single use devices	Catheters, Rinsing	10-730
Single use devices	Catheters, Vascular, Embolectomy/Thrombectomy, Ballon, Venous	10-756
Single use devices	Tubing, Suction	16-779
Single use devices	Strippers, Vein	13-828
Single use devices	Tubes, Bronchial	15-322
Single use devices	Tubes, Connecting	14-188
Single use devices	Valves	14-325
Single use devices	Manifolds	15-587
Single use devices	Catheters, Vascular, Infusion, Central Venous	10-729
	Sterile Procedure Packs acc. §12, MDD	
Single use devices	Casework, General-Purpose	15-896
	Sterile Procedure Packs acc. §12, MDD	

Special terms of validity:

In case of class I products or sterile procedure packs acc. to article 12 (3) of the Directive 93/42/EEC the intervention of ecm is limited to aspects of manufacture concerned with securing and maintaining sterile conditions respectively the conformity with the metrological requirements.

UMDNS Code is optional