



EC Certificate

Production Quality Assurance System Directive 93/42/EEC on Medical Devices (MDD), Annex V (Devices in class I in sterile conditions, sterilised systems or procedure packs)

No. G2S 010393 0531 Rev. 01

Manufacturer CODAN Medizinische Geräte

GmbH & Co KG

Stig Husted-Andersen Strasse 11

23738 Lensahn **GERMANY**

CODAN Medizinische Geräte GmbH & Co KG Facility(ies):

Stig Husted-Andersen Strasse 11, 23738 Lensahn, GERMANY

CODAN PORTUGAL Instrumentos Médicos, S.A.

Rua Stig Husted-Andersen, 4, 2675-492 Odivelas, PORTUGAL

Product Sterile and Non-Sterile Single-Use Medical Category(ies):

Devices for Infusion Therapy as well as

Mixing and Withdraw Products

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for manufacture in accordance with MDD Annex V. This quality assurance system covers those aspects of manufacture concerned with securing and maintaining sterile conditions of the respective devices / device categories and conforms to the requirements of this Directive. It is subject to periodical surveillance. See also notes overleaf.

Report No.: 713169066

Valid from: 2019-12-09

Valid until: 2022-10-03

2019-12-09

Date,

Christoph Dicks Head of Certification/Notified Body

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