

ECCERTIFICATE

Full Quality Assurance System Medical Devices Directive 93/42/EEC Annex II

: Morton Medikal San. ve Tic. A.Ş. Company Name

: İTOB O.S.B. Ekrem Demirtaş Cad. No:9 Tekeli Menderes IZMIR / TURKEY Company Address

: MDD 93/42/EEC Medical Devices Directive - Annex II Related Directives and Annex

(Excluding Section 4)

: Non-sterile Anesthesia and Breathing Circuit - Class Ila Product

> Sterile Bacterial Filter - Class Ila Sterile Catheter Mouth - Class Ila

Sterile Mortonvent Tracheostomy Filter Set - Class Ila Non-sterile Spirometry Filter and Mouth Piece - Class Ila

Sterile Inhalation Holding Chamber - Class Ila Non-sterile Humidifier Chamber - Class Ila

Sterile Extension Line - Class Ila

Sterile Pleural Drainage System - Class Ila

Sterile Y Tur Set - Class Ila

Sterile Yankauer Suction Set - Class Ila

Non-Sterile Disposible Anesthesia Rebreathing Bag - Class Ila

Sterile Video Camera Drape - Class Is Sterile Microscope Drape - Class Is

: 37704, 37706, 37798, 37597, 35795, 37597, 13680, 60699, 12170, 16621, **GMDN**

10817, 46102, 35917, 37450, 12535, 34877

Product Types are attached.

: M.2017.106.8574 Certificate Number

: MD.3375.IB Report Number Initial Assessment Date : 30.05.2017 : 23.06.2017 Registration Date

: 18.08.2017/02 Revision Date /No : 22.06.2022 **Expiry Date**

EXPIRY DATE

UDEM hereby declares that the requirements of Annex II, excluding section 4 of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance audits, defined by Annex II, section 5 of the forementioned directive. According to Annex II, section 4 an EC design- examination certificate is required for placing the Class III devices on the market. This certificate remains as the property of UDEM International Certification Auditing Training Centre Industry and Trade Co. Ltd. to whom if must be returned upon request. The above named company and UDEM must keep a copy of this certificate for 5 years from the registration of the certificate. Usage of the CE mark is under the responsibility of the manufacturer with the completion of EC Declaration of Conformity. The above mentioned company must notify all changes related with the approved product to UDEM. If UDEM will not renew the expiry date of this certificate in question, the mentioned company should stop placing the product on the market. The currency of the certificate can be checked through www.udernital.com.tr.

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Industry

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