

MEDBAR TIBBİ MALZEMELER TURİZM SANAYİ VE TİCARET ANONİM ŞİRKETİ

FATİH MAH. 1142 SOK. NO: 35 SARNIÇ GAZİEMİR – İZMİR – TURKEY

with a scope of

**SURGICAL DRAPE PRODUCTS, IV FLOW REGULATOR PRODUCTS,
KARMAN CANNULA PRODUCTS, ENDOSCOPY MOUTHPIECES, URINE
COLLECTION PRODUCTS, MUCOUS ASPIRATION PRODUCTS,
ARTROCOPY SETS, VOMIT/EMESIS BAG PRODUCTS, SCRUB HAND
BRUSHES, FILTERED MOUTHPIECE PRODUCTS, CERVICAL BRUSH
PRODUCTS, AMNIOTIC POUCH PERFORATORS, FECAL PARASITE
CONCENTRATION PRODUCTS, INTENSIVE CARE PRODUCTS
PROCESSES: PRODUCTION, PACKAGING, STERILIZATION, STORING
DISTRIBUTION AND ETHYLENE OXIDE STERILIZATION SERVICES ARE
UNDER THE SCOPE OF EN ISO 11135 STANDARD**

Medical devices - Quality management systems - Requirements for
regulatory purposes

"Following elements of the standard are excluded"

"7.5.3" "7.5.4" "7.5.9.2"

EN ISO 13485:2016

Certificate No	: M 10326
Initial Certification Date	: 03 October 2019
Certification Date	: 03 October 2019
Expiration Date	: 02 October 2022



Medical Device O.M.S.
TS EN ISO/IEC 17021-1
AB-0006-YS



TÜRKAK BDS NO
YS-694E-E7CB



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Certificate is valid till expiration date, subject to successful completion of periodical surveillance audits.

Please contact above numbers for detailed information.

KIWA BELGELENDİRME HİZMETLERİ
ANONİM ŞİRKETİ

COMPANY HAS BEEN REGISTERED TO OUR
CHAMBER UNDER THE REGISTRATION
NUMBER 365270

PRESENT APPROVAL DOES NOT COVER THE CONTENT OF THE DOCUMENT.
for the SECRETARY GENERAL
OF THE ISTANBUL CHAMBER OF COMMERCE
MUSTAFA TELCİ



Bu belge, tasdik ve imzanın
T.C. TCO'ya ait olduğu tasdik olunur
İşbu tasdik metne şamil değildir
Tasdik No: _____