



# BAYMED



## DECLARATION OF CONFORMITY (UYGUNLUK BEYANI)

**MANUFACTURER: BAYTEKS TEKNİK TEKSTİL SAN. VE TİC. A.Ş**

Organize Sanayi Bölgesi 19 Nolu Cad. No:11/2 MERKEZ/KİLİS

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This declaration of conformity has been prepared in MDR 2017/745 Part II, Article 19, Annex IV; The document is arranged in the home page program for all users.

<b>UDI -DI</b>	: See the Sterile Products table.
<b>PRODUCT</b>	: Sterile Disposable Surgical Gowns, Sterile Disposable Surgical Drapes, Sterile Disposable Surgical Packs.
<b>NOTIFIED BODY</b>	: <b>UDEM Adriatic d.o.o</b> R3 Radnička cesta 54/ R3, Green Gold Centar, IV. kat, 10000, Zagreb, Hırvatistan
<b>ID NO</b>	2696
<b>CND(EMDN) CODE</b>	: <b>See the Sterile Products table.</b>
<b>SRN NUMBER</b>	: TR-MF-000024488
<b>CERTIFICATE NUMBER</b>	: M.2023.MDR.1022
<b>CERTIFICATE VALID UNTIL</b>	: 08.08.2028
<b>CLASSIFICATION</b>	: Class I Sterile (MDR 2017/745 – Annex VIII / Chapter III – Article 4 – Rule 1)
<b>SCOPE</b>	: Sterile Disposable Surgical Gowns, Sterile Disposable Surgical Drapes, Sterile Disposable Surgical Packs.
<b>ADDITIONAL</b>	: EK XI A
<b>APPLIED STANDARDS</b>	: TS EN ISO 13485:2016, ISO 14971:2020, EN ISO 11135:2014, TS EN ISO 15223-1:2021, TS EN ISO 11737-1:2018, EN ISO 11737-2:2020, TS EN ISO 14644-1:2021, TS EN ISO 14644-2:2021, TS EN ISO 10993-1:2021, TS EN ISO 11607-1:2020, TS EN ISO 11607-2:2020, TS EN ISO 11138-1:2017, TS EN ISO 11140-1:2015

The directive for our product is the Council Directive MDR 2017/745 ECC for all versions of medical devices. The Manufacturer of the product, Bayteks Teknik Tekstil San. ve Tic. A.Ş, is responsible for the requirements of this council directive. Our products are not medical devices that contains human blood derivatives, animal products, animal skin, tissues, or blood derivatives or phthalates.

STERILE PRODUCT			
PRODUCT NAME	BASIC UDI	MODEL NO	EMDN
PLAIN DRAPE 80X70 CM	86817441ST30100001445AR	SD-06209-50	T030199



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The products listed in the list above and their contents are classified Class 1 Sterile products. These products, their content, and their accessories do not take part in any other class. We herewith declare that the above mentioned products conforms general requirements of the Council Directive MDR 2017/745 ECC for all versions of Medical Device Directive .

### Applied Directives

Medical Device Directive MDR 2017/745 ECC ANNEX XI A ALL VERSIONS.

**DATE OF ISSUE** : 25.08.2

**024 REV. NO.** : 1

Name and Surname : Yaşar ÇALIŞKAN

Position : Sales Chief

Date of signature : 20.01.2025

Place of Signature : Kilis - Türkiye

