Thoracic Catheter

With Trocar, Standard Tip

- ► Trocar with tri-facetted cutting tip
- ▶ Flexible tubing wall provides leak proof tissue/tubing interface and easy milking
- ▶ Tapered tip for easy and safe insertion
- ▶ Cross side eyes to prevent tissue aspiration
- ▶ Radiopaque line
- ▶ Acccurate positioning depth marks at each cm starting 2 cm from the distal end
- Integral large bore tapered connector



| Ref | Size | Length |
|------------|-------|--------|
| 184 1201 1 | 12 Ch | 225 mm |
| 184 1601 1 | 16 Ch | 235 mm |
| 184 2001 1 | 20 Ch | 390 mm |
| 184 2401 1 | 24 Ch | 390 mm |
| 184 2801 1 | 28 Ch | 390 mm |
| 184 3201 1 | 32 Ch | 390 mm |



Thoracic Catheter

- ▶ Flexible tubing wall provides leak proof tissue/tubing interface
- Atraumatic, rounded, open distal tip
- Easy to milk
- Cross side eyes to prevent tissue aspiration
- ▶ Radiopaque line
- ▶ Accourate positioning depth marks at each cm starting 2 cm from the distal end
- connectors





EG-Zertifikat / EC-Certificate

gem. 93/42/EWG Anhang II ohne (4) / acc. 93/42/EEC Annex II without (4)

Hiermit wird bescheinigt, dass die Firma / This certifies, that the company

Bıçakcılar Tıbbi Cihazlar Sanayi ve Ticaret A.Ş.

Osmangazi Mahallesi, Gazi Caddesi No: 21, Esenyurt 34522 İstanbul Türkiye

für die Produkte / die Kategorie: Liste der Produkte siehe Anlage 1 for the products / product category: List of products see annex 1

Medizinische Einmalartikel und Absauggeräte Disposable medical devices and devices for aspiration and vacuum extraction

ein Qualitätssicherungssystem für die Auslegung, die Fertigung und die Endkontrolle der genannten Produkte nach Maßgabe des Anhang II (ohne Abschnitt 4) der Richtlinie 93/42/EWG anwendet. Zusätzlich zur CE-Kennzeichnung muss die Kennummer der Benannten Stelle angebracht werden. Die Gültigkeit dieses Zertifikats beruht auf der Aufrechterhaltung des Qualitätssicherungssystems in Übereinstimmung mit den Anforderungen der Richtlinie und seiner Überwachung durch die Benannte Stelle gem. Anhang II Abschnitt 5. Das Zertifikat ist unter keinen Umständen übertragbar.

has established a quality system for design, production and final testing acc. to the requirements of Annex II (without section 4) of the directive 93/42/EEC. Additional to the CE-marking the notification number of the Notified Body has to be affixed. The validity of this certificate is based on the maintenance of the quality system in accordance with the requirements of the directive and its surveillance by the Notified Body according Annex II section 5. The certificate may not be transferred under any circumstances.

Reg.-Nr. / Reg.-No. 04 232 980886 Bericht Nr. / Report No. 3524 7139

3524 7139

3526 6290

Zertifizierungsstelle für Medizinprodukte Certification body for medical devices Gültigkeit / Validity von / from 2020-04-16 bis / until 2023-09-16 Edition 8

Essen, 2020-04-16

TÜV NORD CERT GmbH

Langemarckstraße 20

45141 Essen

www.tuev-nord-cert.de

medical@tuev-nord.de





Anlage 1, Blatt 1 von 6 Annex 1, page 1 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse IIb Products of class IIb

Pressure Monitoring Set Leukocyte Filter Set Gamma Leukocyte Filter Set

Produkte der Klasse IIa Products of class IIa

Thoracenthesis Set
Thoracic Catheter
Arterial Needle
Endotracheal Tube
Reinforced Endotracheal Tube
RAE Endotracheal Tube
Nasogastric Catheter
Stomach Catheter
Feeding Catheter
Manifold / Manifold Pressure
Three-Way Stopcock

Bericht Nr. / Report No. 3529 1130

74.78

Zertifizierungsstelle für Medizinprodukte

Certification body for medical devices

TÜV NORD CERT GmbH Langemarckstraße 20

20 45141 Essen

Essen, 2021-05-25

Gültigkeit / Validity

Edition 16

von / from 2021-05-25

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medical@tuev-nord.de





Anlage 1, Blatt 2 von 6 Annex 1, page 2 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse IIa Products of class IIa

Tourniquet Set IV Cannula Suction Catheter Microaggregate Filter Set (Blood Filter Set) Soft Drain Oxygen Catheter Nasal Oxygen Cannula Oxygen Connecting Tube Tracheostomy Tube Extracorporeal PVC Tubing Extracorporeal Tubing Set Quick Prime Set Cardioplegia Set Wound Drainage Set Infusion Pump Set Yankauer Suction Set Suction Connecting Tube Surgical Braided Tape **Nelaton Catheter** Tiemann Catheter

Bericht Nr. / Report No. 3529 1130

7.78

Zertifizierungsstelle für Medizinprodukte Certification body for medical devices Gültigkeit / Validity von / from 2021-05-25 Edition 16

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Anlage 1, Blatt 3 von 6 Annex 1, page 3 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse IIa Products of class IIa

Hydrophilic coated uretheral Catheter IV Filter Set
Aspirators
Blood Transfusion Set
Rectal Catheter
Umbilical Catheter
Angiographic Kit
B-Soft Kit
Aortic Punch
Gas Sampling Line
External Drainage Set
Vent Catheter
Vessel Cannula
Coronary Artery Retraction Clips

Bericht Nr. / Report No. 3529 1130



Zertifizierungsstelle für Medizinprodukte Certification body for medical devices

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Langemarckstraße 20

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von / from 2021-05-25

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Anlage 1, Blatt 4 von 6 Annex 1, page 4 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse Is (steril) Products of class Is (sterile)

Urine Collection Bag Pleural Drainage Set Central Venous Pressure Set **Guedel Airway** Spigot **Extension Lines** Kapkon Connector Straight Connector Straight Luer Connector Y Connector Y Luer Connector Stopper Instopper **Umbilical Cord Clamp** T.U.R. Set / Arthroscopy set Transfer Set Intravenous Infusion Sets Intravenous Infusion Sets / Flowmeter Intravenous Infusion Sets / Burette

Bericht Nr. / Report No. 3529 1130

71.78

Zertifizierungsstelle für Medizinprodukte Certification body for medical devices

Langemarckstraße 20 TÜV NORD CERT GmbH

45141 Essen

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Essen, 2021-05-25

medical@tuev-nord.de

Benannte Stelle Kenn-Nr. 0044 / Notified Body ID. No. 0044

Gültigkeit / Validity von / from 2021-05-25 Edition 16





Anlage 1, Blatt 5 von 6 Annex 1, page 5 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse Is (steril) Products of class Is (sterile)

B-Safe
Intubation Stylet
Combi Stopper
Urimeter
Thoracic Drainage Set
Vaginal Specula
ENEMA Set
I.V. Infusion Set w/B-Flow Flow Regulator
Control Syringe
Meconium Aspiration Connector

Anmerkung: Für Produkte der Klasse I steril beschränkt sich das Zertifizierungsverfahren auf die Aspekte der Herstellungsschritte in Zusammenhang mit der Sterilisation und der Aufrechterhaltung der Sterilität.

Note:

For products of class I sterile the certification process is restricted to the aspects of manufacture concerned with securing and maintaining sterile conditions.

Bericht Nr. / Report No. 3529 1130

74.78

Zertifizierungsstelle für Medizinprodukte Certification body for medical devices Essen, 2021-05-25

Gültigkeit / Validity von / from 2021-05-25

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TÜV NORD CERT GmbH

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45141 Essen

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Anlage 1, Blatt 6 von 6 Annex 1, page 6 of 6

Reg.-Nr. / Reg. No. 04 232 980886

Produkte der Klasse Im (mit Messfunktion) Products of class Im (with measuring function)

Urimeter C.V.P. Set Pleural Drainage Set Volumetric Exerciser (B-Spiro) Infusion Set w/Burette Thoracic Drainage Set

Anmerkung: Für Produkte der Klasse I mit Messfunktion beschränkt sich das Zertifizierungsverfahren auf die

Herstellungsschritte in Zusammenhang mit der Konformität der Produkte mit den messtechnischen

Anforderungen.

For products of class I with measuring functions the certification process is restricted to the aspects of Note:

manufacture concerned with the conformity of the devices with metrological requirements.

Bericht Nr. / Report No. 3529 1130

Zertifizierungsstelle für Medizinprodukte Certification body for medical devices

Essen, 2021-05-25

Gültigkeit / Validity

Edition 16

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TÜV NORD CERT GmbH

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TÜV NORD CERT GmbH, Am TÜV 1, 45307 Essen, Germany

BIÇAKCILAR TIBBI CIHAZLAR SAN. VE TIC. A.Ş. Osmangazi Mahallesi, Gazi Caddesi No: 21, Esenyurt 34522 İstanbul

Turkey

TÜV NORD CERT GmbH

Am TÜV 1 45307 Essen Germany

Phone: +49 201 825 2236

medical@tuev-nord.de tuev-nord-cert.com/en

TÜV®

Reference Contact Direct Dial Date

No.: 8003060047 E-Mail: medical@tuev-nord.de Tel.: +49 201 825 2236 29 June 2023

Notified Body Confirmation Letter Reference: 8003060047

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices

This letter confirms that, TÜV NORD CERT GmbH, a Notified Body (NB) designated against Regulation (EU) 2017/745 (MDR) and identified by the number 0044 on NANDO, has received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the following manufacturer:

BIÇAKCILAR TIBBI CIHAZLAR SAN. VE TIC. A.Ş.

Osmangazi Mahallesi, Gazi Caddesi No: 21,

Esenyurt 34522 İstanbul

Turkey

SRN Number: TR-MF-000022603



Headquarters TÜV NORD CERT GmbH

Am TÜV 1 45307 Essen, Germany

Phone: +49 201 825-0 Fax: +49 201 825-2517 info.tncert@tuev-nord.de tuev-nord-cert.com/en Director
Dipl.-Ing. Wolfgang Wielpütz
Dipl.-Oec. Sandra Gerhartz

Registration Office Amtsgericht Essen HRB 9976 VAT ID No.: DE 811389923 Tax No.: 111/5706/2193 Deutsche Bank AG, Essen BIC (SWIFT-Code): DEUTDEDEXXX IBAN-Code: DE26 3607 0050 0607 8950 00



The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below. Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive. Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but the NB has <u>not</u> yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

In the case of devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively, by the 20 Mar 2023 for the relevant devices.

The transition timelines that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607), are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices excluding Wellestablished technologies (WET - sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition or have a measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

On behalf of the Notified Body,



Digital unterschrieben von Mühlenberg Kevin Datum: 2023.07.05 09:16:27 +02'00'

i. V. Kevin Mühlenberg
 Head of Projectmanagement
 Medical Devices International
 TÜV NORD CERT GmbH
 Notified Body for Medical Devices



Digital unterschrieben von Mestmacher Bodo Datum: 2023.07.05 09:08:26 +02'00'

i. A. Bodo Mestmacher
 Specialist Management
 Medical Devices International
 TÜV NORD CERT GmbH
 Notified Body for Medical Devices

TUVNORD

Table 1: Devices covered by this letter and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

| Device name or Basic UDI-DI (under MDR application) | MDR Device classification (as proposed by the manufacturer and verified at the preapplication stage) | If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device | MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification |
|--|--|--|---|
| Pressure Monitoring Set | Class IIb | N/A | 04232980886 |
| Leukocyte Filter Set | Class IIb | N/A | 04232980886 |
| Gamma Leukocyte Filter Set | Class IIb | N/A | 04232980886 |
| Thoracenthesis Set | Class IIa | N/A | 04232980886 |
| Thoracic Catheter | Class IIa | N/A | 04232980886 |
| Arterial Needle | Class IIa | N/A | 04232980886 |
| Endotracheal Tube | Class IIa | N/A | 04232980886 |
| Reinforced Endotracheal Tube | Class IIa | N/A | 04232980886 |
| RAE Endotracheal Tube | Class IIa | N/A | 04232980886 |
| Nasogastric Catheter | Class IIa | N/A | 04232980886 |
| Stomach Catheter | Class IIa | N/A | 04232980886 |
| Feeding Catheter | Class IIa | N/A | 04232980886 |
| Manifold / Manifold Pressure | Class IIa | N/A | 04232980886 |
| Three -Way Stopcock | Class IIa | N/A | 04232980886 |
| Tourniquet Set | Class IIa | N/A | 04232980886 |
| IV Cannula | Class IIa | N/A | 04232980886 |
| Suction Catheter | Class IIa | N/A | 04232980886 |
| Microaggregate Filter Set (Blood Filter Set) | Class IIa | N/A | 04232980886 |
| Soft Drain | Class IIa | N/A | 04232980886 |
| Oxygen Catheter | Class IIa | N/A | 04232980886 |
| Nasal Oxygen Cannula | Class IIa | N/A | 04232980886 |
| Oxygen Connecting Tube | Class IIa | N/A | 04232980886 |
| Tracheostomy Tube | Class IIa | N/A | 04232980886 |
| Extracorporeal PVC Tubing | Class IIa | N/A | 04232980886 |
| Extracorporeal Tubing Set | Class IIa | N/A | 04232980886 |
| Quick Prime Set | Class IIa | N/A | 04232980886 |
| Cardioplegia Set | Class IIa | N/A | 04232980886 |
| Wound Drainage Set | Class IIa | N/A | 04232980886 |
| Infusion Pump Set | Class IIa | N/A | 04232980886 |
| Yankauer Suction Set | Class IIa | N/A | 04232980886 |
| Suction Connecting Tube | Class IIa | N/A | 04232980886 |
| Surgical Braided Tape | Class IIa | N/A | 04232980886 |
| Nelaton Catheter | Class IIa | N/A | 04232980886 |
| Tiemann Catheter | Class IIa | N/A | 04232980886 |
| Hydrophilic coated uretheral Catheter | Class IIa | N/A | 04232980886 |
| IV Filter Set | Class IIa | N/A | 04232980886 |
| Aspirators | Class IIa | N/A | 04232980886 |
| Blood Transfusion Set | Class IIa | N/A | 04232980886 |
| Rectal Catheter | Class IIa | N/A | 04232980886 |
| Umbilical Catheter | Class IIa | N/A | 04232980886 |
| Angiographic Kit | Class IIa | N/A | 04232980886 |

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| Device name or Basic UDI-DI (under MDR application) | MDR Device classification (as proposed by the manufacturer and verified at the preapplication stage) | If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device | MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification |
|--|--|--|--|
| B -Soft Kit | Class IIa | N/A | 04232980886 |
| Aortic Punch | Class IIa | N/A | 04232980886 |
| Gas Sampling Line | Class IIa | N/A | 04232980886 |
| External Drainage Set | Class IIa | N/A | 04232980886 |
| Vent Catheter | Class IIa | N/A | 04232980886 |
| Vessel Cannula | Class IIa | N/A | 04232980886 |
| Coronary Artery Retraction Clips | Class IIa | N/A | 04232980886 |
| Urine Collection Bag | Class Is | N/A | 04232980886 |
| Pleural Drainage Set | Class Is | N/A | 04232980886 |
| Central Venous Pressure Set | Class Is | N/A | 04232980886 |
| Guedel Airway | Class Is | N/A | 04232980886 |
| Spigot | Class Is | N/A | 04232980886 |
| Extension Lines | Class Is | N/A | 04232980886 |
| Kapkon Connector | Class Is | N/A | 04232980886 |
| Straight Connector | Class Is | N/A | 04232980886 |
| Straight Luer Connector | Class Is | N/A | 04232980886 |
| Y Connector | Class Is | N/A | 04232980886 |
| Y Luer Connector | Class Is | N/A | 04232980886 |
| Stopper | Class Is | N/A | 04232980886 |
| Instopper | Class Is | N/A | 04232980886 |
| Umbilical Cord Clamp | Class Is | N/A | 04232980886 |
| T.U.R. Set /Arthroscopy set | Class Is | N/A | 04232980886 |
| Transfer Set | Class Is | N/A | 04232980886 |
| Intravenous Infusion Sets | Class Is | N/A | 04232980886 |
| Intravenous Infusion Sets / Flowmeter | Class Is | N/A | 04232980886 |
| Intravenous Infusion Sets / Burette | Class Is | N/A | 04232980886 |
| B -Safe | Class Is | N/A | 04232980886 |
| Intubation Stylet | Class Is | N/A | 04232980886 |
| Combi Stopper | Class Is | N/A | 04232980886 |
| Urimeter | Class Is | N/A | 04232980886 |
| Thoracic Drainage Set | Class Is | N/A | 04232980886 |
| Vaginal Specula | Class Is | N/A | 04232980886 |
| ENEMA Set | Class Is | N/A | 04232980886 |
| I.V. Infusion Set w/B-Flow Flow Regulator | Class Is | N/A | 04232980886 |
| Control Syringe | Class Is | N/A | 04232980886 |
| Meconium Aspiration Connector | Class Is | N/A | 04232980886 |
| Urimeter | Class Im | N/A | 04232980886 |
| C.V.P. Set | Class Im | N/A | 04232980886 |
| Pleural Drainage Set | Class Im | N/A | 04232980886 |
| Volumetric Exerciser (B -Spiro) | Class Im | N/A | 04232980886 |
| Infusion Set w/Burette | Class Im | N/A | 04232980886 |
| Thoracic Drainage Set | Class Im | N/A | 04232980886 |



Table 2: Devices covered by this letter and for which the NB is <u>NOT</u> responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

| Device name or Basic UDI-DI (under MDR application) | MDR Device classification (as proposed by the manufacturer and verified at the preapplication stage) | If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device | MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification |
|---|--|--|--|
| N/A | N/A | N/A | N/A |

Confirmation Letter Revision History

| Date | NB internal reference traceable to each version of the letter | Action |
|------------|---|---------------|
| 2023-07-05 | Rev. 0 | Initial issue |
| | | |
| | | |



UYGUNLUK DEKLARASYONU

DECLARATION OF CONFORMITY (Sinif/ Class IIb, IIa, Is, Im)

| Doküman Numarası | DoC-TK2 | Revizyon No: 34 | Tarih: 07,10.2022 | | |
|--|---|---|--------------------------|--|--|
| Document Number | | Revision No | Date | | |
| Üretici Firma Manufacturer | BIÇAKCILAR Tıbbi Cihazlar San. Ve Tic. A.Ş. | | | | |
| Firma adresi | Osmangazi Mahallesi Gazi | Osmangazi Mahallesi Gazi Caddesi No:21 Esenyurt 34522 | | | |
| Manufacturer Address | İSTANBUL/TÜRKİYE | İSTANBUL/TÜRKİYE | | | |
| Onaylanmış Kuruluş & Adresi Notified Body & Address | TÜV NORD CERT GmbH Am TÜV 1 45307 /Essen-G | ermany | | | |

BIÇAKCILAR Tıbbi Cihazlar San. Ve Tic. A.Ş. Yetkili otorite TÜV NORD CERT GmbH (N° 0044) tarafındar değerlendirmiştir.

Bu deklerasyon, Tıbbi Cihaz Direktifi 93/42 EEC Ek VII ve Düzeltme 2007/47/EEC ile uyumlu olarak hazırlanmıştır.

BIÇAKCILAR Tıbbi Cihazlar San. Ve Tic. A.Ş. having been assessed by TÜV NORD CERT GmbH Notified Body

This declaration is made in accordance with Annex VII of the Medical Devices Directive 93/42 EEC and Amendmen 2007/47/EEC

| Uygunluk deklarasyonunda bulunan bütün ürünler için/For all products which are mentioned in the DoC. | | | | | | |
|--|---------------|----------------|---------------------|--|--|--|
| Sertifikalar | Sertifika No | Veriliş Tarihi | Son Kullanma Tarihi | | | |
| Certificates | Certicate No | Date of Issue | Expiry Date | | | |
| EN ISO 13485 (*) | 04 221 980886 | 27.07.2022 | 26.05.2024 | | | |
| 93/42 EEC Ek II / Annex II (4 hariç /without 4) | 04 232 980886 | 16.04.2020 | 16.09.2023 | | | |

(*) EN ISO 13485: 2016 Tıbbi Cihazlar- Kalite Yönetim Sistemleri- Ruhsatlandırma Amaçlı Gereklilikler / Medical Devices-Quality Management Systems- Requirements for Regulatory Purposes

Bıçakcılar Tıbbi Cihazlar A.Ş, Tıbbi Cihazlar Direktifinin 93/42 EEC ve Ek 2007/47/EEC Ek II maddelerine uygun olarak aşağıda belirtilen ürünler için bütün sorumluluğu üstlenir ve ürünün aşağıda belirtilen standardlara ya da diğer düzenleyici mevzuatlara uygunluğunu deklare eder.

Biçakcılar Tibbi Cihazlar A.Ş, Declare under our sole responsibility that the products below to which this declaration relates are in conformity with the following standards or other regulatory laws following the provisions of Medical Device Directive 93/42 EEC and Amendment 2007/47/EEC Annex II.

- ISO 9001: 2015 Kalite Yönetim Sistemleri Gereklilikler/ Quality Management Systems- Requirements
- EN ISO 13485: 2016 Tıbbi Cihazlar- Kalite Yönetim Sistemleri Mevzuat Amaçları Bakımından Şartlar / Medical Devices- Quality Management Systems-Requirements for Regulatory Purposes
- EN ISO 14971 Tıbbi cihazlar Tıbbi cihazlara risk yönetiminin uygulanması / Medical devices Application of risk management to medical devices
- ISO/TR 24971 Tıbbi cihazlar ISO 14971'in uygulanmasına ilişkin kılavuz / Medical devices Guidance on the application of ISO 14971
- EN ISO 10993 Tıbbi cihazların Biyolojik Değerlendirilmesi / Biological Evaluation of Medical Devices
- EN ISO 11135 Sağlık Malzemelerinin Sterilizasyonu-Etilenoksit / Sterilization of Heathcare products-Ethylene oxide
- EN ISO 11607 Son Olarak Steril Edilen Tıbbi Cihazlar için Ambalajlama/ Packaging for terminally sterilized medical device



- EN ISO 11737 Tıbbi Cihazların Sterilizasyonu-Mikrobiyolojik Metodlar /Sterilization of medical devices -Microbiological methods
- ISO 20417 Tıbbi cihazlar İmalatçı tarafından sağlanacak bilgiler / Medical devices Information to be supplied by the manufacturer
- EN ISO 15223 Tibbi cihazlar Tibbi cihaz etiketlerinde, etiketlemede ve sunulacak bilgide kullanılacak semboller / Medical devices Symbols to be used with information to be supplied by the manufacturer
- EN ISO 11138 Sağlık Bakım Ürünlerinin Sterilizasyonu- Biyolojik İndikatörler / Sterilization Of Health Care Products Biological Indicators / Sterilization Of Health Care Products Biological Indicators
- EN ISO 14644 Temiz odalar ve bunlarla ilgili kontrollü ortamlar / Cleanrooms and associated controlled environments
- ISO/TR 20416 Tıbbî cihazlar Üreticiler için Pazar Arz Sonrası Gözetim/ Medical devices Post-market surveillance for manufacturers
- EN 62366-1 Tıbbi cihazlar Bölüm 1: Kullanılabilirlik tekniğinin tıbbi cihazlara uygulanması / Medical devices Part 1: Application of usability engineering to medical devices
- IEC 62366-2 Tıbbi cihazlar Bölüm 2: Kullanılabilirlik tekniğinin tıbbi cihazlara uygulanmasına ilişkin rehberlik / Medical devices Part 2: Guidance on the application of usability engineering to medical devices
- MDCG 2020-6 Eski cihazlar için yeterli klinik kanıt hakkında rehberlik / Guidance on sufficient clinical evidence for legacy devices
- MDCG 2021-25 MDR gerekliliklerinin "eski cihazlara" ve 90/385/EEC veya 93/42/EEC Direktifleri uyarınca 26 Mayıs 2021'den önce piyasaya sürülen cihazlara uygulanması / Application of MDR requirements to "legacy devices" and to devices placed on the market prior to 26 May 2021 in accordance with Directives 90/385/EEC or 93/42/EEC

| | | Sinif lib Ürünle | r i Class Ilb P | roducts | | |
|---------|--|---|-----------------|---------------------|---|---|
| Sıra No | Ürün Referansı | Ürün Adı | GMDN Kodu | Sterilite Durumu | İlgili Ürün Standardı | Ürün Risk Kuralı |
| No | Product Reference | Product Name | GMDN Code | Sterility State | Related Product Standard | Device Risk Rule |
| 1 | 157 00XX 1 015 20XX 1 157 00XX 1G 015 20XX 1G | Lökosit Filtre Seti Leukocyte Filter Set | 35071 | Steril Sterile | ANSI/AAMI BF 64:2012 | Kural 3 / Rule 3 Kural18 / Rule 18 |
| 2 | 040 XXXX 1 400 XXXX 1 | Basınç İzleme Seti Pressure Monitoring Set | 35529 | Steril Sterile | ISO 8536-4: 2019 EN 60601- 1:2006/A1:2013 EN 60601-2-34: 2014 | Kural 10 Rule 10 |

| Sinif Ila Ürünler I Class Ila Products | | | | | | |
|--|--------------------------|--|--------------|---------------------|---|---------------------|
| Sıra No | Ürün Referansı | Ürün Adı | GMDN Kodu | Sterilite Durumu | İlgili Ürün Standardı | Ürün Risk Kuralı |
| No | Product Reference | Product Name | GMDN Code | Sterility State | Related Product Standard | Device Risk Rule |
| 1 | 015 0102 1 | Arteriyal İğne Arterial Needle | 12747 | Steril Sterile | EN ISO 80369-7 (2016) | Kural 6 Rule 6 |
| 2 | 104 1001 1 010 2XXX 1 | Infüzyon pompa seti Infusion pump set | 35833 | Steril Sterile | ISO 8536-4 (2019) EN ISO 8536-8 (2015) | Kural 2 Rule 2 |



| 3 | 113 XXXX 1 114 XXXX 1 | IV Filtre Seti IV Filter Set | 35072 | Steril Sterile | ISO 8536-4 (2019) ISO 80369-7 (2016) | Kural 3 Rule 3 |
|----|--|--|-------|-------------------|---|-------------------|
| 4 | 115 0101 1 | Eksternal Drenaj Büret- 150ml External Drainage Burette- 150ml | 61796 | Steril Sterile | ISO 8536-5 (2004) ISO 20697 (2018) | Kural 2 Rule 2 |
| 5 | 115 0111 1 | Eksternal Drenaj Büret- 150ml Plakalı External Drainage Burette- 150ml - W/plate | 61796 | Steril Sterile | ISO 8536-5 (2004) ISO 20697 (2018) | Kural 2 Rule 2 |
| 6 | 123 1XXX 1 | Üç yollu musluklu uzatma Extention Line w/ Three way Stopcock | 12170 | Steril Sterile | ISO 80369-7 (2016) ISO 8536-9 (2015) | Kural 2 Rule 2 |
| 7 | 145 XXXX 1 146 XXXX 1 014 XXXX 1 095 10XX 1 | B-CAT I.V Kanül B-CAT I.V Cannula | 34905 | Steril Sterile | EN ISO 10555-1 (2013-A1:2018) EN ISO 10555-5 (2013) ISO 80369-7 (2016) | Kural 7 Rule 7 |
| 8 | 150 XXXX 1 151 XXXX 1 154 XXXX 1 155 XXXX 1 015 00XX 1 095 12XX 1 | Kan Transfüzyon Seti Blood Transfusion Set | 38569 | Steril Sterile | ISO 80369-7 (2016) ISO 1135-4 (2015) | Kural 2 Rule 2 |
| 9 | 155 XXXX 1 156 XXXX 1 | Mikroagregat Filtre Seti Microaggregate Filter Set | 35071 | Steril Sterile | ISO 1135-4 (2015) | Kural 3 Rule 3 |
| 10 | 160 XXXX 1 161 XXXX 1 016 XXXX 1 | Yankauer Aspirasyon Ucu Yankauer Suction Handle | 35917 | Steril Sterile | NA | Kural 6 Rule 6 |
| 11 | 162 XXXX 1 | Aspiratör ucu Suction wand | 35917 | Steril Sterile | NA | Kural 6 Rule 6 |
| 12 | 164 XXXX 1 165 XXXX 1 166 XXXX 1 167 XXXX 1 168 XXXX 1 | Yankauer Aspirasyon Seti Yankauer Suction Set | 35917 | Steril Sterile | ISO 20697 (2018) ISO 8836 (2019) | Kural 6 Rule 6 |
| 13 | 168 XXXX X 169 XXXX 1 | Aspirasyon Bağlantı Hortumu Suction Connecting Tube | 16779 | Steril Sterile | ISO 20697 (2018) | Kural 6 Rule 6 |
| 14 | 173 XXXX 1 017 XXXX 1 171 XXXX 1 | B-Vak Doku Drenaj Seti B-Vak Mini Doku Drenaj Seti B-Vak Wound Drainage Set B-Vak Mini Wound Drainage Set | 35824 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
| 15 | 017 11XX 1 | Redon Dren-Trokar Redon Drain-Trochar | 11305 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |



| 16 | 180 XXXX 1 182 XXXX 1 | Toraks Kateteri – Genişleyen Uçlu Thoracic Catheter w/Flared End | 47796 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
|----|--|---|-------|-------------------|---------------------|-------------------|
| 17 | 181 XX01 1 183 XX01 1 | Toraks Kateteri – Tut Çek Konnektörlü Uç Thoracic Catheter w/Pull Through End | 47796 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
| 18 | 184 XXXX 1 | Toraks Kateteri Trokarlı Thoracic Catheter w/Throcar | 47796 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
| 19 | 189 XXXX 1 019 XXXX 1 | Aspirasyon Kateteri (Kapkon Konnektörlü) Suction Catheter (w/Kapkon connector) | 34923 | Steril Sterile | ISO 8836 (2019) | Kural 5 Rule 5 |
| 20 | 190 XXXX 1 191 XX17 1 019 XXXX 1 | Aspirasyon Kateteri Suction Catheter | 34923 | Steril Sterile | ISO 8836 (2019) | Kural 5 Rule 5 |
| 21 | 191 XX11 1 019 XXXX 1 | Aspirasyon Kateteri-Vakum Kontrollü Suction Catheter w/Vacuum Control Connector | 34923 | Steril Sterile | ISO 8836 (2019) | Kural 5 Rule 5 |
| 22 | 019 535X 1 | Aspirasyon Kateteri Vakum Kontrollü Konnektör Kesik Uç Delikli Suction Catheter, w/Vacuum Control Connector Beveled Tip w/Hole | 34923 | Steril Sterile | ISO 8836 (2019) | Kural 5 Rule 5 |
| 23 | 190 XXXX 1 191 XXXX 1 019 XXXX 1 | Aspirasyon Kateteri- Kılıflı Aspirasyon Kateteri- Kılıflı, Eğimli Uç Aspirasyon Kateteri-Vakum Kontrollü Sleeved Suction Catheter Sleeved Suction Catheter, Beveled Tip Suction Catheter w/Vacuum Control Connector | 34923 | Steril Sterile | ISO 8836 (2019) | Kural 5 Rule 5 |
| 24 | 193 XXXX 1 019 XXXX 1 | Mide Kateteri Stomach Catheter | 35415 | Steril Sterile | NA | Kural 5 Rule 5 |
| 25 | 194 XXXX 1 019 XXXX 1 | Nazogastrik Kateter Nasogastric Catheter | 14221 | Steril Sterile | NA | Kural 5 Rule 5 |
| 26 | 195 XX01 1 195 XX05 1 019 XXXX 1 | Nelaton Kateter Nelaton Female Kateter Nelaton Catheter Nelaton Female Catheter | 36125 | Steril Sterile | ISO 20696 (2018) | Kural 5 Rule 5 |
| 27 | 195 XX20 1 019 XXXX 1 | Tiemann Kateteri Tiemann Catheter | 36125 | Steril Sterile | ISO 20696 (2018) | Kural 5 Rule 5 |



| | | B-Soft Hidrofilik Kaplı Kateter | | Cłonii | 100 20000 | Vinal F |
|----|--|--|-------|-------------------|---|-------------------|
| 28 | 196 XXXX 1 | B-Soft Hydrophilic Coated Catheter | 36125 | Steril Sterile | ISO 20696 (2018) | Kural 5 Rule 5 |
| 29 | 196 XX21 1 | B-SOFT Kit | 36125 | Steril Sterile | ISO 20696 (2018) | Kural 5 Rule 5 |
| 30 | 197 XXXX 1 019 XXXX 1 | Beslenme Kateteri Feeding Catheter | 14221 | Steril Sterile | NA | Kural 5 Rule 5 |
| 31 | 197 XX21 1 | Beslenme Kateteri- Enfit Konnektörlü Feeding Catheter- w/ Enfit Connector | 14221 | Steril Sterile | ISO 20695 (2020) | Kural 5 Rule 5 |
| 32 | 198 XXXX 1 019 XXXX 1 | Göbek Kateteri Umbilical Catheter | 10759 | Steril Sterile | ISO 80369-7 (2016) EN ISO 10555-1 (2013-A1:2018) | Kural 7 Rule 7 |
| 33 | 199 XXXX 1 019 XXXX 1 | Rektal Kateter Rectal Catheter | 46202 | Steril Sterile | EN 12439 (1999) | Kural 5 Rule 5 |
| 34 | 300 XXXX 1 304 XXXX 1 310 XXXX 1 311 XXXX 1 312 XXXX 1 315 XXXX 1 776 4001 1 030 XXXX 1 032 XXXX 1 | Ekstrakorporeal Tüp Set Extracorporeal Tubing Set | 35441 | Steril Sterile | ISO 15676 (2016) ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 35 | 305 XXXX X 306 XXXX X 307 XXXX X 030 XXXX 1 | Ekstrakorporeal PVC Hortum Extracorporeal PVC Tubing | 46721 | Steril Sterile | ISO 15676 (2016) | Kural 2 Rule 2 |
| 36 | 320 XXXX 1 032 XXXX 1 | Hızlı Doldurma Seti Quick Prime Set | 35441 | Steril Sterile | ISO 15676 (2016) | Kural 2 Rule 2 |
| 37 | 323 XXXX 1 | Y Adaptör / Perfüzyon Y-Adaptör Y Adapter / Perfusion Y-Adapter | 58824 | Steril Sterile | NA | Kural 2 Rule 2 |
| 38 | 325 XXXX 1 032 XXXX 1 | Kardiopleji Set Cardioplegia Set | 16163 | Steril Sterile | ISO 80369-7 (2016) ISO 1135-4 (2015) | Kural 2 Rule 2 |
| 39 | 330 0XXX 1 | Vent Kateter Vent Catheter | 17613 | Steril Sterile | ISO 20697 (2018) ISO 80369-7 (2016) | Kural 7 Rule 7 |
| 40 | 330 02XX 1 | Vessel Kanül Vessel Cannula | 47798 | Steril Sterile | ISO 80369-7 (2016) | Kural 7 Rule 7 |
| 41 | 330 03XX 1 | Kardiyopleji Adaptörü Cardioplegia Adapter | 58824 | Steril Sterile | NA | Kural 2 Rule 2 |
| 42 | 330 05XX 1 330 0XXX 1 | Turnike set Tourniquet set | 36082 | Steril Sterile | NA | Kural 7 Rule 7 |



| 43 | 332 XXXX 1 | Aortik Punch Aortic Punch | 47914 | Steril Sterile | NA | Kural 6 Rule 6 |
|----|--|---|-------|-------------------|---|-------------------|
| 44 | 135 XXXX 1 138 XXXX 1 340 XXXX 1 341 XXXX 1 | Anjiografik Opak Madde Verme Seti Angiographic Kit | 16545 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 45 | 420 XX01 1 042 000X 1 | Yumuşak Dren Soft Drain | 11305 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
| 46 | 421 0001 1 | Torasentez Seti Thoracentesis Set | 10817 | Steril Sterile | ISO 80369-7 (2016) EN ISO 8669-2 (1996) | Kural 6 Rule 6 |
| 47 | 425 0001 1 042 0001 1 | Göğüs Drenaj Torbası Pleural Drainage Bag | 10817 | Steril Sterile | NA | Kural 7 Rule 7 |
| 48 | 440 4001 1 | Arteriyal Filtre Seti Arterial Filter Set | 33309 | Steril Sterile | NA | Kural 2 Rule 2 |
| 49 | 550 00XX 1 551 00XX 1 055 XXXX 1 | Endotrakeal Tüp (Balonlu/Balonsuz) Endotracheal Tube (Cuffed/Uncuffed) | 46967 | Steril Sterile | EN ISO 5361 (2016) | Kural 5 Rule 5 |
| 50 | 550 8XXX 1 551 8XXX 1 | RAE Endotrakeal Tüp (Balonlu/Balonsuz) RAE Endotracheal Tube (Cuffed/Uncuffed) | 46967 | Steril Sterile | EN ISO 5361 (2016) | Kural 5 Rule 5 |
| 51 | 550 7XXX 1 551 7XXX 1 055 XXXX 1 095 22XX 1 | Spiralli Endotrakeal Tüp (Balonlu/Balonsuz) Reinforced Endotracheal Tube (Cuffed/Uncuffed) | 46569 | Steril Sterile | EN ISO 5361 (2016) | Kural 5 Rule 5 |
| 52 | 551 1XXX 1 | Endotrakeal Tüp (Balonlu, XX mm Stile) Endotracheal Tube (Cuffed with XX mm Stylet) | 46967 | Steril Sterile | EN ISO 5361 (2016) | Kural 5 Rule 5 |
| 53 | 551 20XX 1 | Spiralli Endotrakeal Tüp (Balonlu, XX mm Stile) Reinforced Endotracheal Tube (Cuffed with XX mm Sylet) | 46569 | Steril Sterile | EN ISO 5361 (2016) | Kural 5 Rule 5 |
| 54 | 555 0XXX 1 556 0XXX 1 055 XXXX 1 095 22XX 1 | Trakeostomi Tüp Tracheostomy Tube | 35404 | Steril Sterile | EN 1282-2 (2005- A1:2009) EN ISO 5366 (2016) | Kural 5 Rule 5 |
| 55 | 560 200X 1 560 2001 1 | Nasal Oksijen Kanülü Nasal Oxygen Cannula | 35201 | Steril Sterile | NA | Kural 2 Rule 2 |
| 56 | 563 XXXX 1 056 XXXX 1 | Oksijen Kateteri Oxygen Catheter | 35203 | Steril Sterile | NA | Kural 2 Rule 2 |



| 57 | 565 XXXX 1 056 XXXX 1 | Oksijen Bağlantı Hortumu Oxygen Connecting Tube | 12875 | Steril Sterile | EN 1617 (1997) ISO 20697 (2018) | Kural 2 Rule 2 |
|----|--|---|-------|-------------------|--|-------------------|
| 58 | 573 0X7X 1 057 0X7X 1 | Gaz Örnekleme Hattı Gas Sampling Line | 45566 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 59 | 723 XX70 1 726 XX70 1 724 XXXX 1 072 XXXX 1 | Cerrahi Örme Bant Surgical Braided Tape | 36082 | Steril Sterile | NA | Kural 7 Rule 7 |
| 60 | 760 XXXX 1 076 XXXX 1 | Üç Yollu Musluk Three Way Stopcock | 32172 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 61 | 765 XXXX 1 076 XXXX 1 | Manifold <i>Manifold</i> | 32172 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 62 | 790 XX01 1 079 XXXX 1 | Redon Dren Redon Drain | 11305 | Steril Sterile | ISO 20697 (2018) | Kural 7 Rule 7 |
| 63 | 330 0450 1 | Koroner Arter Retraksiyon Klipsi- 3.0mm Coronary Artery Retraction Clips-3.0mm | 47991 | Steril Sterile | NA | Kural 6 Rule 6 |
| 64 | 330 0451 1 | Koroner Arter Retraksiyon Klipsi- 5.0mm Coronary Artery Retraction Clip- 5.0mm | 47991 | Steril Sterile | NA | Kural 6 Rule 6 |

| | Sinif Im Ürünler / Class Im Products | | | | | | |
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| Sıra No | Ürün Referansı | Ürün Adı | GMDN Kodu | Sterilite Durumu | İlgili Ürün Standardı | Ürün Risk Kuralı | |
| No | Product Reference | Product Name | GMDN Code | Sterility State | Related Product Standard | Device Risk Rule | |
| 1 | 186 XXXX 2 | B-Spiro Nefes Egzersiz Cihazı B-Spiro Volumetric Exerciser | 31266 | Non-Steril Non-Sterile | NA | Kural 5 Rule 5 | |

| | Sinif Is Ürünler / Class Is Products | | | | | | |
|------------|--|---------------------------------------|--------------|---------------------|---|---------------------|--|
| Sıra No | Ürün Referansı | Ürün Adı | GMDN Kodu | Sterilite Durumu | İlgili Ürün Standardı | Ürün Risk Kuralı | |
| No | Product Reference | Product Name | GMDN Code | Sterility State | Related Product Standard | Device Risk Rule | |
| 1 | 100 XXXX 1 101 XXXX 1 102 XXXX 1 103 XXXX 1 010 XXXX 1 | I.V. İnfüzyon Seti I. V. Infusion Set | 58977 | Steril Sterile | ISO 8536-4 (2019) EN ISO 8536-8 (2015) | Kural 2 Rule 2 | |



| 2 | 106 XXXX 1 107 000X 1 | Damla Ayar Seti Flow Regulator | 36244 | Steril Sterile | ISO 80369-7 (2016) ISO 8536-4 (2019) | Kural 2 Rule 2 |
|----|--|--|----------------|-------------------|--|-------------------|
| 3 | 106 000X 1 107 000X 1 010 05XX 1 | Infüzyon Seti-Damla Ayarlı I.V. Infusion Set w/Flowmeter | 58977 | Steril Sterile | ISO 8536-4 (2019) ISO 8536-8 (2015) | Kural 2 Rule 2 |
| 4 | 120 XXXX 1 121 XXXX 1 122 XXXX 1 012 XXXX 1 | Uzatma Hatları Extention Lines | 12170 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 5 | 125 0005 1 125 0001 1 012 XXXX 1 | Stoper / İnstoper Stopper/ Instopper | 31667 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 6 | 125 0007 1 012 XXXX 1 | Kombi Stoper Combi stopper | 31667 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 7 | 125 0010 1 012 XXXX 1 | Transfer Set | 41222 | Steril Sterile | NA | Kural 1 Rule 1 |
| 8 | 125 10XX 1 130 XXXX 1 131 XXXX 1 012 XXXX 1 | B Safe | 42727 | Steril Sterile | ISO 80369-1 (2018) ISO 80369-7 (2016) ISO 80369-20 (2015) | Kural 2 Rule 2 |
| 9 | 131 00XX 1 132 00XX 1 133 XXXX 1 124 XXXX 1 013 XXXX 1 | B Safe Valfli Uzatma- İkili/Üçlü/T-Konnektörlü Extension Line w/B-Safe Duo/Triple/T-Connector | 12170 | Steril Sterile | ISO 80369-1 (2018) ISO 80369-7 (2016) ISO 80369-20 (2015) | Kural 2 Rule 2 |
| 10 | 135 XXXX 1 138 XXXX 1 013 80XX 1 | Basınca Dayanıklı Uzatma Hatları Pressure Extention Lines | 35529 | Steril Sterile | ISO 80369-7 (2016) ISO 8536-9 (2015) | Kural 2 Rule 2 |
| 11 | 222 XXXX 1 223 XXXX 1 226 XXXX 1 022 XXXX 1 | İdrar Torbası Urine Collection Bag | 58921 58922 | Steril Sterile | EN ISO 8669-2 (1996) | Kural 1 Rule 1 |
| 12 | 022 XXXX 1 | Bacak İdrar Torbası Leg Bag | 58924 | Steril Sterile | NA | Kural 1 Rule 1 |
| 13 | 228 XXXX 1 022 XXXX 1 | Lavman Seti Lavman Torba Enema Set Enema Bag | 35050 | Steril Sterile | NA | Kural 5 Rule 5 |
| 14 | 230 0001 1 023 0001 1 | Göbek Kordon Klempi Umbilical Cord Clamp | 43998 | Steril Sterile | TS 6782: 1989 (T1:1994) | Kural 1 Rule 1 |
| 15 | 235 0001 1 023 0001 1 | Konik Konnektör Conical Connector | 44545 | Steril Sterile | NA | Kural 1 Rule 1 |



| 16 | 236 XXXX 1 | Hortum Konnektörü | 44545 | Steril | N/A | Kural 2 |
|----|--|---|-------|-------------------|--------------------------|-------------------|
| 10 | 023 0001 1 | Tubing Connector | 44040 | Sterile | NA | Rule 2 |
| 17 | 236 1001 1 | Mekonyum Aspiratör Konnektörü Meconium Aspirator Connector | 35917 | Steril Sterile | NA | Kural 2 Rule 2 |
| 18 | 238 0001 1 238 0011 1 023 XXXX 1 | Kateter Tıkacı Spigot | 31667 | Steril Sterile | NA | Kural 1 Rule 1 |
| 19 | 240 0001 1 024 0001 1 | Kapkon Konnektör Kapkon Connector | 44545 | Steril Sterile | NA | Kural 1 Rule 1 |
| 20 | 430 XXXX 1 043 XXX1 1 | TUR Set | 46102 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 21 | 450 XXX1 1 045 XXXX 1 | Artroskopi Set Arthroscopy Set | 46102 | Steril Sterile | ISO 80369-7 (2016) | Kural 2 Rule 2 |
| 22 | 550 0001 1 550 0002 1 550 0003 1 055 XXXX 1 | Entübasyon Stilet Entubation Stylet | 37469 | Steril Sterile | NA | Kural 5 Rule 5 |
| 23 | 595 10XX 1 | Vajinal Spekulum Vaginal Specula | 37468 | Steril Sterile | TS 5537:1988 T3: 2003 | Kural 5 Rule 5 |
| 24 | 750 XXXX 1 075 XXXX 1 | Düz Konnektör Straight Connector | 35338 | Steril Sterile | NA | Kural 2 Rule 2 |
| 25 | 751 XXXX 1 075 XXXX 1 | Düz Luer Konnektör Straight Luer Connector | 35338 | Steril Sterile | NA | Kural 2 Rule 2 |
| 26 | 754 XXXX 1 075 XXXX 1 | Y Konnektör Y Connector | 35338 | Steril Sterile | NA | Kural 2 Rule 2 |
| 27 | 755 XXXX 1 075 XXXX 1 | Y Luer Konnektör Y Luer Connector | 35338 | Steril Sterile | NA | Kural 2 Rule 2 |
| 28 | 900 XXXX 1 095 90XX 1 090 XXXX 1 | Guedel Havayolu Guedel Airway | 42424 | Steril Sterile | EN ISO 5364 (2016) | Kural 2 Rule 2 |
| 29 | 034 XXXX 1 | Kontrol Şırıngası Control Syringe | 15286 | Steril Sterile | ISO 80369-7 -(2016) | Kural 2 Rule 2 |

| | Sinif Is-Im Ürünler / Class is & Im Products | | | | | | |
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| Sıra No | Ürün Referansı | Ürün Adı | GMDN Kodu | Sterilite Durumu | İlgili Ürün Standardı | Ürün Risk Kuralı | |
| No | Product Reference | Product Name | GMDN Code | Sterility State | Related Product Standard | Device Risk Rule | |
| 1 | 010 XXXX 1 105 XXXX 1 095 11XX 1 | I.V. Infüzyon Seti-Büretli I.V. Infusion Set-w/Burette | 12159 | Steril Sterile | EN ISO 8536-5 (2013) ISO 80369-7 (2016) | Kural 2 Rule 2 | |



| 2 | 011 XXXX 1 110 0001 1 | C. V. P. SET Central Venous Pressure Monitoring Set | 35529 | Steril Sterile | ISO 8536-4 (2019) ISO 80369-7 (2016) | Kural 2 Rule 2 |
|---|--------------------------|---|-------|-------------------|---|-------------------|
| 3 | 017 XXXX 1 175 XXXX 1 | BPDS- Göğüs drenaj seti Pleural Drainage Set | 10817 | Steril Sterile | ISO 20697 (2018) | Kural 1 Rule 1 |
| 4 | 017 XXXX 1 176 200X 1 | BTDS –Toraks drenaj seti Thoracic Drainage Set | 10817 | Steril Sterile | ISO 20697 (2018) | Kural 1 Rule 1 |
| 5 | 227 XXXX 1 022 XXXX 1 | Ürimetre <i>Urimeter</i> | 32072 | Steril Sterile | EN ISO 8669-2 (1996) | Kural 1 Rule 1 |
| 6 | 022 7XXX 1 227 10XX 1 | Urimetre İdrar Torbalı Urimeter w/Urine Bag | 32072 | Steril Sterile | EN ISO 8669-2 (1996) | Kural 1 Rule 1 |
| 7 | 027 1023 1 | Ürimetre 500 Plus - İğnesiz Num. Portlu-Çek Valf Urimeter 500 Plus- Needleless Sample-Check Valve | 32072 | Steril Sterile | EN ISO 8669-2 (1996) | Kural 1 Rule 1 |
| 8 | 022 7404 1 | Urimeter 500 Plus Safety | 32072 | Steril Sterile | EN ISO 8669-2 (1996) | Kural 1 Rule 1 |

Açıklama: XXXX ürünün farklı uzunluk, ölçü gibi farklılıklarını ifade etmektedir. Explanation:XXXX means different length, sizes etc. product.
NA: İlgili ürün standardı bulunmamaktadır./ There is no related product standard.

| ONAY / APPROVAL | |
|---|---|
| Yayın Yeri ve İmza Tarihi Signature Date and Place of Issue | TURKEY/ 07.10.2022 |
| Yetkili kişinin adı, ünvar Name, title, signature of authori | nı, imzası ve firma kaşesi zed person with company cachet |
| Kalite Güvence Uzmanı Quality Assurance Specialist | Kalite ve Regülasyon Yöneticisi Quailty and Regulatory Exercutive |
| Selda ÇAKMAK | Aysel YILDIRIM |
| Seida ÇAKMAK Kalite Güvence Uzmanı Quality Assurance Specialist | Ayzel YILDIRIM Kalite ve Regülasyon Yöneticisi Quality and Regulatory Executive |