

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

BIÇAKCILAR TIBBİ CİHAZLAR SAN. VE TIC. A.Ş.
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TÜV®

Reference

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Date

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:

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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 not 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(AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or provided evidence that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 Well-established 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
unterscriben 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

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 pre- application 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
Thoracentesis 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 urethral 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

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre- application 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 NOT 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 pre-application 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

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,
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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 Kennnummer 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
3526 6208
3526 6290



Gültigkeit / Validity
von / from 2020-04-16
bis / until 2023-09-16
Edition 8

Zertifizierungsstelle für Medizinprodukte
Certification body for medical devices

Essen, 2020-04-16

TÜV NORD CERT GmbH Langemarckstraße 20 45141 Essen www.tuev-nord-cert.de medical@tuev-nord.de

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



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Zentralstelle der Länder
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ANLAGE / ANNEX

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

Thoracentesis 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



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

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ANLAGE / ANNEX

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

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ANLAGE / ANNEX

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 urethral 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



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ANLAGE / ANNEX

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



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Edition 16

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ANLAGE / ANNEX

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

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ANLAGE / ANNEX

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.

Note: *For products of class I with measuring functions the certification process is restricted to the aspects of manufacture concerned with the conformity of the devices with metrological requirements.*

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