



AT-OS Srl | Viale del Lavoro, 19 - 37030 Colognola ai Colli - Verona - ITALY
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Partita IVA: IT 02719270239

DECLARATION OF CONFORMITY

Seller : TURKUAZ BIYOMEDIKAL TEKN. VE SAG. HIZM. SAN. TIC. LTD. STI.
KAZIM OZALP MAH. HAFTA SOK. NO: 23/2 06610 CANKAYA / ANKARA /TURKEY

Manufacturer : AT-OS S.r.l.
Viale del Lavoro, 19 Colognola ai Colli (VR) – Italy

We, as manufacturer of this product, herewith declare under our sole responsibility that the below mentioned products designed especially for the seller mentioned above:

Product : Machine for washing and disinfection

UMBNS Code : 17671

Model Code : PLUSHER WDC 8, PLUSHER WDS 12, PLUSHER WDS18

Classification : IIb (According to MDD, Annex IX)

meet the provisions of the following EC Council Directives and Standards. All supporting documentation's are retained under the premises of the manufacturer and the notified body.

Directives : Medical Device Directive – COUNCIL DIRECTIVE of 14 June 1993 concerning medical devices (MDD 93/42/EEC), Annex I; 2007/47/EEC: updating to Medical Device Directive 93/42/EEC.

Standards : Standards applicable to this product are EN ISO 13485:2012; DIN EN 1717:2001; DIN EN 13077:2004; EN 61010-1:2010; EN 61010-2-040:2005; EN 61326-1:2006 + Recommendation NB-MED/2.12 Rec 1; EN ISO 15883-1:2009; EN ISO 15883-2:2009.

Notified Body : TÜV SÜD Product Service GmbH.
Ridler Str. 65, D-80339 München, Germany

Conf. Asses. Route : Annex II.3.


AT-OS S.r.l.
Francesco Avesani
(Regulatory Office)

06.09.2017



DECLARATION OF CONFORMITY

Seller : TURKUAZ BIYOMEDİKAL TEKN. VE SAĞ. HİZM. SAN. TİC. LTD. ŞTİ.
KAZIM OZALP MAH. HAFTA SOK. NO: 23/2 06610 CANKAYA / ANKARA /TURKEY

Manufacturer : AT-OS S.r.l.
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Notified Body : TÜV SÜD Product Service GmbH.
Ridler Str. 65, D-80339 München, Germany

Conf. Asses. Route : Annex II.3.

TURKUAZ BIYOMEDİKAL TEKNOLOJİLER
VE SAĞ. HİZ. SAN. TİC. LTD. ŞTİ.
Kazım Özalp Mah. Hafta Sok. No: 23/2
G.O.P. - Cankaya / ANKARA / TURKEY
Tel: +90 850 840 8 828 Fax: +90 312 911 25 28
Cumhurbaşkanlığı MDD 47/10624583
Mersis No: 0671052153300016

M. Hilmi MİRELİ
CEO
06.09.2017



Add value.
Inspire trust.

TÜV SÜD Product Service GmbH · Ridlerstr. 65 · 80339 Munich · Germany

AT-OS S.r.l.
Viale del Lavoro 19
37030 COGNOLA AI COLLI (VR)
ITALY

Your reference/letter of	Our reference/name	Tel. extension/Email	Fax extension	Date	Page
041151	713294212; ITA1992645_CL; ITA200220003828_CL; ITA200220006001; 713349046	+39 3489010334 Paolo.Bolelli@tuvsud.com	-	2024-10-07	1 of 25

TÜV SÜD Product Service GmbH
Confirmation Letter
CL 041151 0015 Rev. 02

Reference: 713294212 | ITA1992645_CL | ITA200220003828_CL | ITA200220006001 | 713349046

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 (in the following referenced as MDR) as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices.

With this letter TÜV SÜD Product Service GmbH, designated under MDR and identified by the number 0123 on NANDO, confirms that we have 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 above stated manufacturer with the following SRN Number:

SRN Number: **IT-MF-000028623**

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 TÜV SÜD Product Service GmbH 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 TÜV SÜD Product Service GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

Registered Office: Munich
Trade Register Munich HRB 85 742
UniCredit Bank AG · BIC HYVEDEMMXXX
IBAN DE13 7002 0270 0048 8522 11
VAT ID No. DE129484267
Information pursuant to § 2 [1] DL-InfoV
(Germany) at tuvsud.com/imprint

Supervisory Board:
Holger Lindner (Chairman)
Board of Management:
Walter Reithmaier (CEO)
Patrick van Welij

TÜV SÜD Product Service GmbH
Certification Body for Medical Products
Ridlerstr. 65
80339 Munich
Germany

tuvsud.com/ps
Hotline: +49 89 50084-747





If 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.

The transition timelines in accordance Article 120 (3) of MDR 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, 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 (except 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, 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)

The issuance of the first confirmation letter is free of charge. We reserve the right to invoice further copies, amendments and / or changes of the confirmation letter according to effort.

For confirmation letter validity see www.tuvsud.com/ps-cert?q=CL_041151_0015

In case of inquiries please contact medical_devices@tuvsud.com.

On behalf of the Notified Body TÜV SÜD Product Service GmbH,
2024-10-07

TÜV SÜD Product Service GmbH
Medical and Health Services

TÜV SÜD Product Service GmbH
Medical and Health Services

Paolo Bolelli
[Paolo Bolelli \(Oct 7, 2024 13:58 GMT+2\)](#)

Paolo Bolelli
Conformity Assessment Responsible (CARE)

Michael Mauermeir
[Michael Mauermeir \(Oct 7, 2024 12:58 GMT+2\)](#)

Michael Mauermeir
Application Reviewer



Table 1: Devices covered by this letter and for which TÜV SÜD Product Service GmbH 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 during application review)	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
AF2.45MEG-P17 AF2.45MEG-P21 AF2.45MEG-P23 AF2.45MEG-P19 AF2.45MEG-P11 AF2.45MEG-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45MEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45MEV-P17 AF2.45MEV-P21 AF2.45MEV-P23 AF2.45MEV-P19 AF2.45MEV-P11 AF2.45MEV-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45MEV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45MG-P17 AF2.45MG-P21 AF2.45MG-P23 AF2.45MG-P19 AF2.45MG-P11 AF2.45MG-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45MG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45MV-P17 AF2.45MV-P21 AF2.45MV-P23 AF2.45MV-P19 AF2.45MV-P11 AF2.45MV-P06 BASIC UDI-DI:	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number:	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
805673649AF2LK	☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	AF2.45MV	
AF2.45PEG-P17 AF2.45PEG-P21 AF2.45PEG-P23 AF2.45PEG-P19 AF2.45PEG-P11 AF2.45PEG-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45PEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45PEV-P17 AF2.45PEV-P21 AF2.45PEV-P23 AF2.45PEV-P19 AF2.45PEV-P11 AF2.45PEV-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45PEV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45PG-P17 AF2.45PG-P21 AF2.45PG-P23 AF2.45PG-P19 AF2.45PG-P11 AF2.45PG-P06 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.45PG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.45PV-P17 AF2.45PV-P21 AF2.45PV-P23 AF2.45PV-P19 AF2.45PV-P11 AF2.45PV-P06	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
BASIC UDI-DI: 805673649AF2LK	☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	Individual Article number: AF2.45PV	
AF2.60MEG-P17 AF2.60MEG-P21 AF2.60MEG-P23 AF2.60MEG-P19 AF2.60MEG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60MEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60METG-P17 AF2.60METG-P21 AF2.60METG-P23 AF2.60METG-P19 AF2.60METG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60METG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60METV-P17 AF2.60METV-P21 AF2.60METV-P23 AF2.60METV-P19 AF2.60METV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60METV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60MEV-P17 AF2.60MEV-P21 AF2.60MEV-P23 AF2.60MEV-P19 AF2.60MEV-P11	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted)	☐ N/A or	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
BASIC UDI-DI: 805673649AF2LK	☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60MEV	
AF2.60MG-P17 AF2.60MG-P21 AF2.60MG-P23 AF2.60MG-P19 AF2.60MG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60MG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60MV-P17 AF2.60MV-P21 AF2.60MV-P23 AF2.60MV-P19 AF2.60MV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60MV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60PEG-P17 AF2.60PEG-P21 AF2.60PEG-P23 AF2.60PEG-P19 AF2.60PEG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60PEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60PETG-P17 AF2.60PETG-P21 AF2.60PETG-P23 AF2.60PETG-P19	☐ Class III ☐ Class IIb implantable (non-exempted)	☐ N/A or	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AF2.60PETG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60PETG	
AF2.60PETV-P17 AF2.60PETV-P21 AF2.60PETV-P23 AF2.60PETV-P19 AF2.60PETV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60PETV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60PEV-P17 AF2.60PEV-P21 AF2.60PEV-P23 AF2.60PEV-P19 AF2.60PEV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.60PEV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
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AF2.60PV-P17 AF2.60PV-P21	☐ Class III	☐ N/A	☒ Certification as follows: G1 041151 0014 Rev. 00;



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AF2.60XMEG-P17 AF2.60XMEG-P21 AF2.60XMEG-P23 AF2.60XMEG-P19 AF2.60XMEG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60MEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
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AF2.60XMG-P17 AF2.60XMG-P21 AF2.60XMG-P23 AF2.60XMG-P19 AF2.60XMG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60MG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60XMV-P17 AF2.60XMV-P21 AF2.60XMV-P23 AF2.60XMV-P19 AF2.60XMV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60MV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60XPEG-P17 AF2.60XPEG-P21 AF2.60XPEG-P23 AF2.60XPEG-P19 AF2.60XPEG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60PEG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60XPEV-P17 AF2.60XPEV-P21 AF2.60XPEV-P23 AF2.60XPEV-P19 AF2.60XPEV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60PEV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
	☐ Class III implantable custom-made-device		
AF2.60XPG-P17 AF2.60XPG-P21 AF2.60XPG-P23 AF2.60XPG-P19 AF2.60XPG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60PG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.60XPV-P17 AF2.60XPV-P21 AF2.60XPV-P23 AF2.60XPV-P19 AF2.60XPV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X60PV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90AG-P17 AF2.90AG-P21 AF2.90AG-P23 AF2.90AG-P19 AF2.90AG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90AG/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90ATG-P17 AF2.90ATG-P21 AF2.90ATG-P23 AF2.90ATG-P19 AF2.90ATG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90ATG/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
	☐ Class I devices with measuring function ☐ Class III implantable custom-made-device		
AF2.90ATV-P17 AF2.90ATV-P21 AF2.90ATV-P23 AF2.90ATV-P19 AF2.90ATV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90ATV/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90AV-P17 AF2.90AV-P21 AF2.90AV-P23 AF2.90AV-P19 AF2.90AV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90AV/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90BAG-P17 AF2.90BAG-P21 AF2.90BAG-P23 AF2.90BAG-P19 AF2.90BAG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90BAG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90BATG-P17 AF2.90BATG-P21 AF2.90BATG-P23 AF2.90BATG-P19 AF2.90BATG-P11 BASIC UDI-DI:	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
805673649AF2LK	☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	Individual Article number: AF2.90BATG	
AF2.90BATV-P17 AF2.90BATV-P21 AF2.90BATV-P23 AF2.90BATV-P19 AF2.90BATV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90BATV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90BAV-P17 AF2.90BAV-P21 AF2.90BAV-P23 AF2.90BAV-P19 AF2.90BAV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90BAV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90BG-P17 AF2.90BG-P21 AF2.90BG-P23 AF2.90BG-P19 AF2.90BG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90BG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90BV-P17 AF2.90BV-P21 AF2.90BV-P23 AF2.90BV-P19 AF2.90BV-P11	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted)	☐ N/A or	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
BASIC UDI-DI: 805673649AF2LK	☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90BV	
AF2.90G-P17 AF2.90G-P21 AF2.90G-P23 AF2.90G-P19 AF2.90G-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90G/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90V-P17 AF2.90V-P21 AF2.90V-P23 AF2.90V-P19 AF2.90V-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.90V/860	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90XBAG-P17 AF2.90XBAG-P21 AF2.90XBAG-P23 AF2.90XBAG-P19 AF2.90XBAG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X90BAG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90XBAG-P17 AF2.90XBAG-P21 AF2.90XBAG-P23 AF2.90XBAG-P19	☐ Class III ☐ Class IIb implantable (non-exempted)	☐ N/A or	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AF2.90XBAG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X90BAG	
AF2.90XBAV-P17 AF2.90XBAV-P21 AF2.90XBAV-P23 AF2.90XBAV-P19 AF2.90XBAV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X90BAV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90XBG-P17 AF2.90XBG-P21 AF2.90XBG-P23 AF2.90XBG-P19 AF2.90XBG-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X90BG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.90XBV-P17 AF2.90XBV-P21 AF2.90XBV-P23 AF2.90XBV-P19 AF2.90XBV-P11 BASIC UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.X90BV	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-2-P11 AWD655-2-P10	☐ Class III	☐ N/A	☒ Certification as follows: G1 041151 0014 Rev. 00;



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AWD655-2X-P11 AWD655-2X-P10 AWD655-2E-P11 AWD655-2E-P10 AWD655-2XE-P11 AWD655-2XE-P10 BASIC UDI-DI: 805673649AWD655FC	☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-2	NB# 0123
AWD655-2H-P11 AWD655-2H-P10 AWD655-2HX-P11 AWD655-2HX-P10 AWD655-2HE-P11 AWD655-2HE-P10 AWD655-2HXE-P11 AWD655-2HXE-P10 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-2H	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-8-P01 AWD655-8-P02 AWD655-8-P03 AWD655-8-P04 AWD655-8-P05 AWD655-8-P06 AWD655-8-P07 AWD655-8-P08 AWD655-8-P09 AWD655-8XE-P01 AWD655-8XE-P02 AWD655-8XE-P03 AWD655-8XE-P04 AWD655-8XE-P05 AWD655-8XE-P06 AWD655-8XE-P07 AWD655-8XE-P08 AWD655-8XE-P09 AWD655-8E-P01 AWD655-8E-P02 AWD655-8E-P03 AWD655-8E-P04 AWD655-8E-P05 AWD655-8E-P06	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-8	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classifica- tion (as proposed by the manufacturer and verified during appli- cation review)	If the MDR device is a sub- stitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
AWD655-8E-P07 AWD655-8E-P08 AWD655-8E-P09 AWD655-8X-P01 AWD655-8X-P02 AWD655-8X-P03 AWD655-8X-P04 AWD655-8X-P05 AWD655-8X-P06 AWD655-8X-P07 AWD655-8X-P08 AWD655-8X-P09 BASIC UDI-DI: 805673649AWD655FC			
AWD655-8L-P01 AWD655-8L-P02 AWD655-8L-P03 AWD655-8L-P04 AWD655-8L-P05 AWD655-8L-P06 AWD655-8L-P07 AWD655-8L-P08 AWD655-8L-P09 AWD655-8XEL-P01 AWD655-8XEL-P02 AWD655-8XEL-P03 AWD655-8XEL-P04 AWD655-8XEL-P05 AWD655-8XEL-P06 AWD655-8XEL-P07 AWD655-8XEL-P08 AWD655-8XEL-P09 AWD655-8EL-P01 AWD655-8EL-P02 AWD655-8EL-P03 AWD655-8EL-P04 AWD655-8EL-P05 AWD655-8EL-P06 AWD655-8EL-P07 AWD655-8EL-P08 AWD655-8EL-P09 AWD655-8XL-P01 AWD655-8XL-P02 AWD655-8XL-P03 AWD655-8XL-P04	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corre- sponding device under MDD/AIMDD Individual Article number: AWD655-8L	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AWD655-8XL-P05 AWD655-8XL-P06 AWD655-8XL-P07 AWD655-8XL-P08 AWD655-8XL-P09 BASIC UDI-DI: 805673649AWD655FC			
AWD655-8L-SC-P01 AWD655-8L-SC-P02 AWD655-8L-SC-P03 AWD655-8L-SC-P04 AWD655-8L-SC-P05 AWD655-8L-SC-P06 AWD655-8L-SC-P07 AWD655-8L-SC-P08 AWD655-8L-SC-P09 AWD655-8EL-SC-P01 AWD655-8EL-SC-P02 AWD655-8EL-SC-P03 AWD655-8EL-SC-P04 AWD655-8EL-SC-P05 AWD655-8EL-SC-P06 AWD655-8EL-SC-P07 AWD655-8EL-SC-P08 AWD655-8EL-SC-P09 AWD655-8XL-SC-P01 AWD655-8XL-SC-P02 AWD655-8XL-SC-P03 AWD655-8XL-SC-P04 AWD655-8XL-SC-P05 AWD655-8XL-SC-P06 AWD655-8XL-SC-P07 AWD655-8XL-SC-P08 AWD655-8XL-SC-P09 AWD655-8XEL-SC-P01 AWD655-8XEL-SC-P02 AWD655-8XEL-SC-P03 AWD655-8XEL-SC-P04 AWD655-8XEL-SC-P05 AWD655-8XEL-SC-P06 AWD655-8XEL-SC-P07 AWD655-8XEL-SC-P08 AWD655-8XEL-SC-P09 BASIC UDI-DI:	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-8L-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
805673649AWD655FC			
AWD655-10-P01 AWD655-10-P02 AWD655-10-P14 AWD655-10-P15 AWD655-10-P16 AWD655-10X-P01 AWD655-10X-P02 AWD655-10X-P14 AWD655-10X-P15 AWD655-10X-P16 AWD655-10E-P01 AWD655-10E-P02 AWD655-10E-P14 AWD655-10E-P15 AWD655-10E-P16 AWD655-10XE-P01 AWD655-10XE-P02 AWD655-10XE-P14 AWD655-10XE-P15 AWD655-10XE-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-10-SC-P01 AWD655-10-SC-P02 AWD655-10-SC-P14 AWD655-10-SC-P15 AWD655-10-SC-P16 AWD655-10X-SC-P01 AWD655-10X-SC-P02 AWD655-10X-SC-P14 AWD655-10X-SC-P15 AWD655-10X-SC-P16 AWD655-10E-SC-P01 AWD655-10E-SC-P02 AWD655-10E-SC-P14 AWD655-10E-SC-P15 AWD655-10E-SC-P16 AWD655-10XE-SC-P01 AWD655-10XE-SC-P02 AWD655-10XE-SC-P14 AWD655-10XE-SC-P15 AWD655-10XE-SC-P16 BASIC UDI-DI:	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
805673649AWD655FC			
AWD655-10A-P01 AWD655-10A-P02 AWD655-10A-P14 AWD655-10A-P15 AWD655-10A-P16 AWD655-10AE-P01 AWD655-10AE-P02 AWD655-10AE-P14 AWD655-10AE-P15 AWD655-10AE-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10A	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-10A-SC-P01 AWD655-10A-SC-P02 AWD655-10A-SC-P14 AWD655-10A-SC-P15 AWD655-10A-SC-P16 AWD655-10AE-SC-P01 AWD655-10AE-SC-P02 AWD655-10AE-SC-P14 AWD655-10AE-SC-P15 AWD655-10AE-SC-P16 AWD655-10A-2SC-P01 AWD655-10A-2SC-P02 AWD655-10A-2SC-P14 AWD655-10A-2SC-P15 AWD655-10A-2SC-P16 AWD655-10AE-2SC-P01 AWD655-10AE-2SC-P02 AWD655-10AE-2SC-P14 AWD655-10AE-2SC-P15 AWD655-10AE-2SC-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10A-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-10AD-P01 AWD655-10AD-P02 AWD655-10AD-P14 AWD655-10AD-P15 AWD655-10AD-P16 AWD655-10ADE-P01 AWD655-10ADE-P02	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AWD655-10ADE-P14 AWD655-10ADE-P15 AWD655-10ADE-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	Individual Article number: AWD655-10AD	
AWD655-10AD-SC-P01 AWD655-10AD-SC-P02 AWD655-10AD-SC-P14 AWD655-10AD-SC-P15 AWD655-10AD-SC-P16 AWD655-10ADE-SC-P01 AWD655-10ADE-SC-P02 AWD655-10ADE-SC-P14 AWD655-10ADE-SC-P15 AWD655-10ADE-SC-P16 AWD655-10AD-2SC-P01 AWD655-10AD-2SC-P02 AWD655-10AD-2SC-P14 AWD655-10AD-2SC-P15 AWD655-10AD-2SC-P16 AWD655-10ADE-2SC-P01 AWD655-10ADE-2SC-P02 AWD655-10ADE-2SC-P14 AWD655-10ADE-2SC-P15 AWD655-10ADE-2SC-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10AD-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-10D-P01 AWD655-10D-P02 AWD655-10D-P14 AWD655-10D-P15 AWD655-10D-P16 AWD655-10DX-P01 AWD655-10DX-P02 AWD655-10DX-P14 AWD655-10DX-P15 AWD655-10DX-P16 AWD655-10DE-P01 AWD655-10DE-P02 AWD655-10DE-P14 AWD655-10DE-P15 AWD655-10DE-P16 AWD655-10DXE-P01	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10D	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AWD655-10DXE-P02 AWD655-10DXE-P14 AWD655-10DXE-P15 AWD655-10DXE-P16 BASIC UDI-DI: 805673649AWD655FC			
AWD655-10D-SC-P01 AWD655-10D-SC-P02 AWD655-10D-SC-P14 AWD655-10D-SC-P15 AWD655-10D-SC-P16 AWD655-10DX-SC-P01 AWD655-10DX-SC-P02 AWD655-10DX-SC-P14 AWD655-10DX-SC-P15 AWD655-10DX-SC-P16 AWD655-10DE-SC-P01 AWD655-10DE-SC-P02 AWD655-10DE-SC-P14 AWD655-10DE-SC-P15 AWD655-10DE-SC-P16 AWD655-10DXE-SC-P01 AWD655-10DXE-SC-P02 AWD655-10DXE-SC-P14 AWD655-10DXE-SC-P15 AWD655-10DXE-SC-P16 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-10D-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-15A-P17 AWD655-15AE-P17 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-15A	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-15A-SC-P17 AWD655-15AE-SC-P17 AWD655-15A-2SC-P17	☐ Class III ☐ Class IIb implantable (non-exempted)	☐ N/A or	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
AWD655-15AE-2SC-P17 AWD655-15A-3SC-P17 AWD655-15AE-3SC-P17 BASIC UDI-DI: 805673649AWD655FC	☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-15A-SC	
AWD655-15AD-P17 AWD655-15ADE-P17 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-15AD	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AWD655-15AD-SC-P17 AWD655-15ADE-SC-P17 AWD655-15AD-2SC-P17 AWD655-15ADE-2SC-P17 AWD655-15AD-3SC-P17 AWD655-15ADE-3SC-P17 BASIC UDI-DI: 805673649AWD655FC	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AWD655-15AD-SC	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
AF2.686SG AF2.686SAG AF2.686SV AF2.686SAV AF2.45SG Basic UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☐ N/A or ☒ Identification of the corresponding device under MDD/AIMDD Individual Article number: AF2.686SG AF2.686SAG AF2.686SV AF2.686SAV AF2.45SG	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	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
REVO REVO-PRO Basic UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☐ Class IIb / Class IIb implantable (exempted) ☒ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ N/A or ☐ Identification of the corresponding device under MDD/AIMDD Individual Article number:	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123
TIVA10-1M TIVA10-1M-1HS TIVA10-1M-1HS-E TIVA10-1M-1HS-X TIVA10-1M-1HS-XE TIVA10-1M-2HS TIVA10-1M-2HS-E TIVA10-1M-2HS-X TIVA10-1M-2HS-XE TIVA10-1M-E TIVA10-1M-X TIVA10-1M-XE TIVA10-1V TIVA10-1V-1HS TIVA10-1V-1HS-E TIVA10-1V-2HS TIVA10-1V-2HS-E TIVA10-1V-E TIVA10-2M TIVA10-2M-1HS TIVA10-2M-1HS-E TIVA10-2M-1HS-X TIVA10-2M-1HS-XE TIVA10-2M-2HS TIVA10-2M-2HS-E TIVA10-2M-2HS-X TIVA10-2M-2HS-XE TIVA10-2M-E TIVA10-2M-X TIVA10-2M-XE TIVA10-2V TIVA10-2V-1HS TIVA10-2V-1HS-E TIVA10-2V-2HS TIVA10-2V-2HS-E	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ N/A or ☐ Identification of the corresponding device under MDD/AIMDD Individual Article number:	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classifica- tion (as proposed by the manufacturer and verified during appli- cation review)	If the MDR device is a sub- stitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
TIVA10-2V-E TIVA15-1V TIVA15-1V-HS TIVA15-2V TIVA15-2V-HS TIVA2 TIVA2-E TIVA2-H TIVA2-HE TIVA2-HX TIVA2-HXE TIVA2-X TIVA2-XE TIVA8-1M TIVA8-E-1M TIVA8-WD-HS-1M TIVA8-WD-1M TIVA8-WD-E-1M TIVA8-WD-HS-E-1M TIVA8-WD-HS-X-1M TIVA8-WD-HS-XE-1M TIVA8-WD-X-1M TIVA8-WD-XE-1M TIVA8-X-1M TIVA8-XE-1M Basic UDI-DI: 805673649AF2LK			
SWD.8.1MD SWD.8.1MD.L SWD.8.1MD.SC.L SWD.12.1MD SWD.12.2MD SWD.12.1VD SWD.12.2VD SWD.12.1MD.SC SWD.12.2MD.SC SWD.12.1VD.SC SWD.12.2VD.SC SWD.18.1VD SWD.18.2VD SWD.18.1VD.SC SWD.18.2VD.SC Basic UDI-DI: 805673649AF2LK	☐ Class III ☐ Class IIb implantable (non-exempted) ☒ Class IIb / Class IIb implantable (exempted) ☐ Class IIa ☐ Class I devices in sterile condition ☐ Class I devices with measuring function ☐ Class III implantable custom-made-device	☒ N/A or ☐ Identification of the corre- sponding device under MDD/AIMDD Individual Article number:	☒ Certification as follows: G1 041151 0014 Rev. 00; NB# 0123



Table 2: Devices covered by this letter and for which TÜV SÜD Product Service GmbH 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 during application review)	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
Not applicable	☒ N/A	☒ N/A	☒ N/A

Confirmation Letter Version History

Date	TÜV SÜD Product Service GmbH internal reference traceable to each version of the letter	Action
2024-03-19	713294212; ITA1992645_CL	Initial issue
2024-06-24	ITA200220003828_CL	Addition of device (Basic UDI-DI 805673649AF2LK); removal of double entry on page 4
2024-10-07	ITA200220006001; 713349046	Addition of device families "REVO", "TIVA", and "SWD" (re-branding of already covered devices); Correction of classification of devices with Basic UDI-DI "805673649AWD655FC" – corrected from class IIa to class IIb



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
ZLG-BS-244.10.08



Product Service

EC Certificate

Full Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)

(Devices in Class IIa, IIb or III)

No. G1 041151 0014 Rev. 00

Manufacturer:

AT-OS S.r.l.

Viale del Lavoro 19

37030 Colognola ai Colli (VR)

ITALY

Facility(ies):

AT-OS S.r.l.

Viale del Lavoro 19, 37030 Colognola ai Colli (VR), ITALY

**Product Category(ies): Washer-disinfectors for
non active medical devices**

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

Report No.:

ITA1339913

Valid from:

2019-08-27

Valid until:

2024-05-26

Date,

2019-08-27

Stefan Preiß

Head of Certification/Notified Body



GOGREEN CERTIFICATE 2023

TURKUAZ BIYOMEDIKAL TEK.SAG.HIZ.SAN.TIC.LT.ST

offset a total of **219.06 kg CO₂e**
for 2023 with GoGreen Products and Services.

DHL Group has offset the greenhouse gas emissions generated by transportation and logistics through worldwide, registered climate protection projects.

More details about the DHL GoGreen Projects Portfolio and selection criteria can be found at:
<https://group.dhl.com/climate-protection-projects>

Michiel Greeven
Executive Vice President
Global Commercial DHL Express

This certificate is issued by DHL Group. The greenhouse gas emissions stated on this certificate (reported as CO₂e*) include emissions from transport and logistics as well as upstream emissions from fuel and energy production. The emissions have been calculated and offset via expenditures on climate protection projects as mentioned above. SGS (Société Générale de Surveillance) has verified the tracked greenhouse gas emissions and the related offsets against the Carbon Management System and according to the „Greenhouse Gas Protocol - Product Life Cycle Accounting and Reporting Standard” for the period of 01.01.2023 to 31.12.2023.

* CO₂e: The CO₂ equivalents for emission compensation include carbon dioxide (CO₂) as well as further GHG emissions such as methane (CH₄).





CERTIFICATE

TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA / ANKARA / TÜRKİYE

Has been independently audited and found to comply with the requirements of the following standard.

Bağımsız olarak denetlenmiş ve aşağıdaki standardın gerekliliklerine uygun olduğu görülmüştür.

ISO 10002:2018

The Customer Satisfaction Management System is applicable to:

Müşteri Memnuniyeti Yönetim Sistemi:

This certificate is given within the scope of the below:

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS& DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

Bu sertifika aşağıdaki kapsam dahilinde verilmiştir.

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

Certificate Number : D44027N

Certificate Initial Issue Date : 10.01.2024

Certificate Issue Date : 09.01.2025

Certificate Validity Date : 08.01.2026

Certification Period : 3 Years

Revision Number : 00

Sertifika No

Sertifika İlk Düzenleme Tarihi

Sertifika Düzenleme Tarihi

Sertifika Geçerlilik Tarihi

Belgelendirme Periyodu

Revizyon No

Certificate validity information www.dcsertification.com you can check at. DCS CERTIFICATION INSPECTION PRIVATE COMPANY D-17, Daliganj, Lucknow, Uttar Pradesh 226020, India Web: www.dcsertification.com • E-Mail: info@dcsertification.com



CERTIFICATION MANAGER



CERTIFICATE

TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.

KAZIM ÖZALP MAH. HAFTA SOK. NO:23/2
ÇANKAYA / ANKARA / TÜRKİYE

*Has been assessed and found to comply with the requirements of:
Denetlenmiş ve aşağıdaki standardın gerekliliklerine uygunluğu görülmüştür:*

ISO 13485:2016

*Medical Devices-Quality Management System is applicable to:
Tıbbi Cihazlar Kalite Yönetim Sistemi*

DESIGN, DEVELOPMENT, MANUFACTURE, SALES AND SERVICE OF STEAM STERILISERS, PLASMA STERILISERS AND CONSUMABLES, LTSF STERILISERS, AUTOCLAVES, WASHING AND DISINFECTION DEVICES, ULTRASONIC WASHERS, BAG SEALING DEVICES, ETHYLENE OXIDE STERILISERS, HOSPITAL STAINLESS EQUIPMENT, CONTAINER SYSTEMS AND ACCESSORIES, STERILISATION ROLLS AND BAGS, AUTOCLAVE TAPES, CREPE PAPER, DISINFECTANTS AND DETERGENTS FOR THE MEDICAL SECTOR, SALES OF WASHING AND STERILISATION CONTROL PRODUCTS, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS

MEDİKAL SEKTÖRE YÖNELİK BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, STERİLİZASYON RULOLARI VE POŞETLERİNİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN SATIŞI

Certificate Number: 2025/MDQMS/000896
Belge Numarası: 2025/MDQMS/000896

Initial Certification Date: 14.03.2025
İlk Belgelendirme Tarihi: 14.03.2025

Certification Period: 3 Years
Belgelendirme Periyodu: 3 Yıl

Certificate Validity Date: 13.03.2026
Belge Geçerlilik Tarihi: 13.03.2026

IQR Sertifikasyon Onayı



IQR ULUSLARARASI BELGELENDİRME HİZMETLERİ LTD.ŞTİ.

Beşevler Mah. Kocayunus Sk. No:3 Arslan Han Plaza K:2 Nilüfer / BURSA

Tel.: +90.224.266 00 16 Faks: +90.224.249 41 13 www.iqrcert.com e-posta: info@iqrcert.com



Sertifika

CERTIFICATE

TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA/ANKARA/TÜRKİYE

ISO 14001:2015

Kapsam/Scope

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS & DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

IAF KOD:7-12-14-17-18-19-23

Bu sertifika ile yukarıda adı geçen kuruluşun Çevre Yönetim Sistemi gerekliliklerini karşıladığı tasdik olunur.

This is to certify that the above mentioned company meets the requirement of Environmental Management System.

Sertifika No / Certification Number	: MTS-10676
İlk Kayıt Tarihi / Date of Initial Reg.	: 5.03.2018
Basım Tarihi / Date of Certificate	: 25.12.2024
Geçerlilik Tarihi / Date of Expiry	: 4.03.2026
Revizyon / Revision	: 007

Bu sertifika kuruluşun belgelendirme şartlarına uyması ve yılda en az 1 kez yapılacak olan gözetim denetimlerinin başarılı geçmesi halinde üç yıllık sertifikasyon periyodu bitiş tarihine kadar geçerlidir.

This certificate is valid until the end of the three-year certification period if the organization complies with the certification requirements and the surveillance audits to be carried out at least once a year are completed successfully.



Genel Müdür / General Manager

SİGMACERT ULUSLARARASI BELGELENDİRME EĞİTİM VE TEST HİZMETLERİ LTD. ŞTİ.

● Bahçekapı Mahallesi Sanayi Bulvarı Şaşmaz Business Plaza No: 18/25 Etimesgut/Ankara

● www.sigmacert.com.tr ● belgelendirme@sigmacert.com.tr ☎ +90 312 385 08 85

YS.FR.129 Rev.03 18.09.2024



CERTIFICATE

TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA / ANKARA / TÜRKİYE

Has been independently audited and found to comply with the requirements of the following standard.

Bağımsız olarak denetlenmiş ve aşağıdaki standardın gerekliliklerine uygun olduğu görülmüştür.

ISO / IEC 27001:2022

The Information Security Management System is applicable to:

Bilgi Güvenliği Yönetim Sistemi:

This certificate is given within the scope of the below:

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS& DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

Bu sertifika aşağıdaki kapsam dahilinde verilmiştir.

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

Applicability Statement / Uygulanabilirlik Bildirgesi: SOA.00 Rev 00/04.06.2024

Certificate Number	: D44025W	Sertifika No
Certificate Initial Issue Date	: 10.01.2024	Sertifika İlk Düzenleme Tarihi
Certificate Issue Date	: 09.01.2025	Sertifika Düzenleme Tarihi
Certificate Validity Date	: 08.01.2026	Sertifika Geçerlilik Tarihi
Certification Period	: 3 Years	Belgelendirme Periyodu

Revision Number

00

Revizyon No

Certificate validity information www.dcs-certification.com you can check at. DCS CERTIFICATION INSPECTION PRIVATE COMPANY D-17, Daliganj, Lucknow, Uttar Pradesh 226020, India Web: www.dcs-certification.com • E-Mail: info@dcs-certification.com



CERTIFICATION MANAGER

Signature



Şertifika

CERTIFICATE

**TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE
SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.**

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA/ANKARA/TÜRKİYE

ISO 45001:2018

Kapsam/Scope

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS & DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

IAF KOD:7-12-14-17-18-19-23

Bu sertifika ile yukarıda adı geçen kuruluşun İş Sağlığı ve Güvenliği Yönetim Sistemi gerekliliklerini karşıladığı tasdik olunur.

This is to certify that the above mentioned company meets the requirement of Occupational Health & Safety Management System.

Sertifika No / Certification Number	: MTS-42480
İlk Kayıt Tarihi / Date of Initial Reg.	: 25.12.2024
Basım Tarihi / Date of Certificate	: 25.12.2024
Geçerlilik Tarihi / Date of Expiry	: 24.12.2025
Revizyon / Revision	: 000

Bu sertifika kuruluşun belgelendirme şartlarına uyması ve yılda en az 1 kez yapılacak olan gözetim denetimlerinin başarılı geçmesi halinde üç yıllık sertifikasyon periyodu bitiş tarihine kadar geçerlidir.

This certificate is valid until the end of the three-year certification period if the organization complies with the certification requirements and the surveillance audits to be carried out at least once a year are completed successfully.



Genel Müdür / General Manager

SIGMACERT ULUSLARARASI BELGELENDİRME EĞİTİM VE TEST HİZMETLERİ LTD. ŞTİ.

● Bahçekapı Mahallesi Sanayi Bulvarı Şaşmaz Business Plaza No: 18/25 Etimesgut/Ankara

● www.sigmacert.com.tr ● belgelendirme@sigmacert.com.tr ● +90 312 385 08 85

YS.FR.129 Rev.03 18.09.2024



CERTIFICATE

TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA / ANKARA / TÜRKİYE

Has been independently audited and found to comply with the requirements of the following standard.

Bağımsız olarak denetlenmiş ve aşağıdaki standardın gerekliliklerine uygun olduğu görülmüştür.

ISO 50001:2018

The Energy Management System is applicable to:

Enerji Yönetim Sistemi:

This certificate is given within the scope of the below:

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS& DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

Bu sertifika aşağıdaki kapsam dahilinde verilmiştir.

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

Certificate Number	: D44026W	Sertifika No
Certificate Initial Issue Date	: 10.01.2024	Sertifika İlk Düzenleme Tarihi
Certificate Issue Date	: 09.01.2025	Sertifika Düzenleme Tarihi
Certificate Validity Date	: 08.01.2026	Sertifika Geçerlilik Tarihi
Certification Period	: 3 Years	Belgelendirme Periyodu
Revision Number	: 00	Revizyon No

Certificate validity information www.dcsertification.com you can check at. DCS CERTIFICATION INSPECTION PRIVATE COMPANY D-17, Daliganj, Lucknow, Uttar Pradesh 226020, India Web: www.dcsertification.com • E-Mail: info@dcsertification.com



CERTIFICATION MANAGER



Sertifika

CERTIFICATE

**TURKUAZ BİYOMEDİKAL TEKNOLOJİLER VE
SAĞLIK HİZMETLERİ SAN. TİC. LTD. ŞTİ.**

KAZIM ÖZALP MAH. HAFTA SOK. NO: 23/2 ÇANKAYA/ANKARA/TÜRKİYE

ISO 9001:2015

Kapsam/Scope

BUHAR STERİLİZATÖRLERİNİN, PLAZMA STERİLİZATÖRLERİ VE SARFLARININ, LTSF STERİLİZATÖRLERİNİN, OTOKLAVLARIN, YIKAMA VE DEZENFEKSİYON CİHAZLARININ, ULTRASONİK YIKAYICILARIN, POŞET KAPAMA CİHAZLARININ, ETİLEN OKSİT STERİLİZATÖRLERİNİN, HASTANE PASLANMAZ EKİPMANLARININ, KONTEYNER SİSTEMLERİ VE AKSESUARLARININ, YIKAMA VE STERİLİZASYON KONTROL ÜRÜNLERİNİN, STERİLİZASYON RULOLARI VE POŞETLERİNİN, BOWIE-DICK TEST PAKETLERİNİN, KİMYASAL İNDİKATÖRLERİN, BİYOLOJİK İNDİKATÖRLERİN, OTOKLAV BANTLARININ, KREPE KAĞITLARININ VE DEZENFEKTAN VE DETERJANLARIN DİZAYNI, GELİŞTİRMESİ, ÜRETİMİ, SATIŞI VE SERVİSİ

DESIGN, DEVELOPMENT, PRODUCTION, SALES AND SERVICE OF STEAM STERILIZERS, PLASMA STERILIZERS & DISPOSABLES, LTSF STERILIZERS, AUTOCLAVES, WASHER DISINFECTORS, ULTRASONIC CLEANERS, SEALERS, ETHYLENE OXIDE STERILIZERS, STAINLESS HOSPITAL EQUIPMENTS, CONTAINER SYSTEMS & ACCESSORIES, STERILIZATION & WASHING CONTROL PRODUCTS, STERILIZATION ROLLS AND POUCHES, BOWIE-DICK TEST PACKAGES, CHEMICAL INDICATORS, BIOLOGICAL INDICATORS, AUTOCLAVE TAPES, WRAPPING PAPERS AND DISINFECTANTS & DETERGENTS

IAF KOD:7-12-14-17-18-19-23

Bu sertifika ile yukarıda adı geçen kuruluşun Kalite Yönetim Sistemi gerekliliklerini karşıladığı tasdik olunur.

This is to certify that the above mentioned company meets the requirement of Quality Management System.

Sertifika No / Certification Number	: MTS-10675
İlk Kayıt Tarihi / Date of Initial Reg.	: 5.03.2018
Basım Tarihi / Date of Certificate	: 25.12.2024
Geçerlilik Tarihi / Date of Expiry	: 4.03.2026
Revizyon / Revision	: 007

Bu sertifika kuruluşun belgelendirme şartlarına uyması ve yılda en az 1 kez yapılacak olan gözetim denetimlerinin başarılı geçmesi halinde üç yıllık sertifikasyon periyodu bitiş tarihine kadar geçerlidir.

This certificate is valid until the end of the three-year certification period if the organization complies with the certification requirements and the surveillance audits to be carried out at least once a year are completed successfully.



Genel Müdür / General Manager

SIGMACERT ULUSLARARASI BELGELENDİRME EĞİTİM VE TEST HİZMETLERİ LTD. ŞTİ.

📍 Bahçekapı Mahallesi Sanayi Bulvarı Şaşmaz Business Plaza No: 18/25 Etimesgut/Ankara

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