

REPUBLICA



MOLDOVA

# CERTIFICAT DE ÎNREGISTRARE

**SOCIETATEA CU RĂSPUNDERE LIMITATĂ "TEHNOMEDICA"**  
ESTE ÎNREGISTRATĂ LA CAMERA ÎNREGISTRĂRII DE STAT

*Numărul de indentificare de stat - codul fiscal*

**1002600053256**

*Data înregistrării*

**17.04.2002**

*Data eliberării*

**16.02.2005**

**Bolboceanu Adela, registruator de stat**

*Funcția, numele, prenumele persoanei  
care a eliberat certificatul*

*semnătura*

**MD 0027040**



Nr. CIF26-842.2020  
Data: 13 Februarie 2020

**CERTIFICAT  
PRIVIND EXISTENTA CONTURILOR CURENTE**

Prin prezentul, **Mobiasbanca - OTP Group S.A.**, codul băncii (BIC): **MOBBMD22**, confirmă că compania **TEHNOMEDICA S.R.L.** cod fiscal (IDNO) **1002600053256**, detine următoarele conturi curente la Mobiasbanca - OTP Group S.A., Sucursala. 26 Negruzzi:

1. **MDL - MD65MO2224ASV98310887100**
2. **EUR - MD06MO2224ASV98311097100**

  
L.S.  
Numele, Prenumele si Semnătura  
Director sucursalei „Gheorghe Mocanu”



Executor :Eduard Cilic  
Tel: 022-812-150



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 010066 0426 Rev. 00**

**Manufacturer:** **AESCULAP AG**  
Am Aesculap-Platz  
78532 Tuttlingen  
GERMANY

**Product Category(ies):** **Implants, Instruments and Devices**  
**(for detailed information see attachment)**

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.:** 713159626

**Valid from:** 2019-07-27

**Valid until:** 2024-05-26

**Date,** 2019-07-16

Stefan Preiß  
Head of Certification/Notified Body

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 ZERTIFIKAT ◆ CERTIFICATE ◆ 認證證書 ◆ CERTIFICADO ◆ CERTIFICAT



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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 010066 0426 Rev. 00**

**Facility(ies):**

AESULAP AG  
Am Aesculap-Platz, 78532 Tuttlingen, GERMANY

- Surgical and dental instruments
- Joint implants (hip, knee)
- Spinal implants
- Implants for osteosynthesis
- Neurosurgical vascular implants
- Products for ligature
- Motor systems
- High frequency surgery devices
- Endoscopic systems
- Navigation system
- Surgical suction pumps
- Implants for replacement of connective tissue
- Vascular prostheses and accessories  
and other surgical accessories
- Collagen implants

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 ZERTIFIKAT ◆ CERTIFICATE ◆ 認證證書 ◆ CERTIFICADO ◆ CERTIFICAT

**Declaration**

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The certification body of TÜV Süd Management Service GmbH and the TÜV Süd Product Service GmbH confirm that we,

**AESULAP AG  
AM AESULAP-PLATZ  
78532 TUTTLINGEN / GERMANY**

have established and are maintaining a quality management system according to

**ISO 9001:2015**

(Certificate Registration No.: 12 100 21724 TMS)

**EN ISO 13485:2016**

(Certificate No.: Q5 17 03 10066 408)

for the following area

**Development, Production and Distribution of Implants, Instruments, Containers,  
Devices, Suture Material, Tissue Adhesives and Procedure Kits.**

Furthermore we have implemented the conformity assessment procedure as per annex II, clause 3 of the Medical Device Directive 93/42/EEC of June 14<sup>th</sup>, 1993 for medical products.

By labeling the products

**Aesculap Product Groups  
as per attached list**

with the CE mark

we, **AESULAP AG** confirm,  
that we follow the essential requirements  
according to MDD 93/42/EEC Annex I.

TUTTLINGEN, 2018-03-13

AESULAP AG

i. V.



Thomas Marquard  
Regulatory Affairs

i. A.



Denise Hermle  
Regulatory Affairs

<b>Aesculap Product Groups</b>
Surgical, diagnostic and dental instruments
Joint Implants (Hip, Knee)
Spinal Implants
Implants for osteosynthesis
Neurosurgical Vascular Implants
Products for Ligature
Motor Systems
Sterilization Containers and Accessoires
High Frequency Surgery Devices
Endoscopic Systems
Navigation Systems
Surgical Suction Pumps
Special Suture-Sets
Implants for Replacement of Connective Tissue
Tissue Adhesives
Vascular Prosthesis and Accessories
Local Haemostatics
Other Surgical Accessories



# Certificate

No. Q5 010066 0435 Rev. 00

**Holder of Certificate:** **AESCULAP AG**  
Am Aesculap-Platz  
78532 Tuttlingen  
GERMANY

**Certification Mark:**



**Scope of Certificate:** **Design and development, production, technical service and distribution of implants, instruments, instrument management systems, containers, devices, tissue adhesives**

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). See also notes overleaf.

**Report No.:** 713175266

**Valid from:** 2020-06-01  
**Valid until:** 2023-05-31

**Date,** 2020-05-27

Christoph Dicks  
Head of Certification/Notified Body

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# Certificate

No. Q5 010066 0435 Rev. 00

**Applied Standard(s):** EN ISO 13485:2016  
Medical devices - Quality management systems -  
Requirements for regulatory purposes  
(ISO 13485:2016)  
DIN EN ISO 13485:2016

**Facility(ies):** AESCULAP AG  
Am Aesculap-Platz, 78532 Tuttlingen, GERMANY  
  
AESCULAP AG  
Carl-Braun-Str. 1, 34212 Melsungen, GERMANY

- Surgical and dental instruments
- Joint implants (hip, knee)
- Spinal implants
- Implants for osteosynthesis
- Neurosurgical vascular implants
- Products for ligature
- Motor systems
- Sterilization containers and accessories
- High frequency surgery devices
- Endoscopic systems
- Navigation systems
- Surgical suction pumps
- Implants for replacement of connective tissue
- Tissue adhesives
- Vascular prostheses and accessories
- Local haemostatics
- Other surgical accessories
- Collagen implants





Management Service

# CERTIFICATE

The Certification Body  
of TÜV SÜD Management Service GmbH

certifies that

## Aesculap AG

Am Aesculap-Platz, 78532 Tuttlingen, Germany  
Carl-Braun-Straße 1, 34212 Melsungen, Germany

has established and applies  
a Quality Management System for

**Design and Development, Technical Service, Production and Distribution of  
Implants, Instruments, Containers, Devices,  
Suture Material and Tissue Adhesive**

### Aesculap AG Tuttlingen

- Surgical and dental instruments
- Joint Implants (hip, knee)
- Spinal Implants
- Implants for Osteosynthesis
- Neurosurgical Vascular Implants
- Motor systems
- Sterilization containers and accessories
- High frequency surgery devices
- Endoscopic systems
- Navigation systems
- Surgical suction pumps
- Veterinary instrumentation
- Other surgical accessories
- Instrument Management System
- Collagen implants

### Aesculap AG Melsungen

- Implants for replacement of connective tissue
- Tissue adhesive
- Local haemostatic

An audit was performed, Order No. **70062209**.

Proof has been furnished that the requirements according to

## ISO 9001:2015

are fulfilled.

The certificate is valid from **2020-06-01** until **2023-05-31**.

Certificate Registration No.: **12 100 21724 TMS**.

Product Compliance Management  
Munich, 2020-05-20



CERTIFICAT



CERTIFICADO



СЕРТИФИКАТ



認證證書



CERTIFICATE



ZERTIFIKAT