



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
BS-MDR-099



EU Production Quality Assurance Certificate

Regulation (EU) 2017/745 on Medical Devices, Annex XI Part A

Certificate No. G26 057924 0020 Rev. 00

Manufacturer:

Ningbo Advan Electrical Co., Ltd.

Industrial Development Zone
Fuhai Town
315332 Cixi City, Ningbo, Zhejiang Province
PEOPLE'S REPUBLIC OF CHINA

SRN Manufacturer - CN-MF-000018731

Authorized Representative:

CMC Medical Devices & Drugs S.L.
C/Horacio Lengo Nº 18, CP 29006 Málaga, SPAIN

The quality management system has been evaluated in accordance with Regulation (EU) 2017/745, Annex XI Part A with a positive result.

Details on devices covered by the quality management system are described on the following page(s). The report referenced below summarises the results of the assessment and includes reference to relevant CS, harmonised standards and test reports.

The certified quality management system is subject to periodical surveillance.

If class I devices in sterile conditions, with measuring function, or reusable surgical instruments are covered by this certificate, the audit was limited to the respective aspects relating to

- establishing, securing, and maintaining sterile conditions,
- conformity of the devices with the metrological requirements,
- reuse of the device, in particular cleaning, disinfection, sterilization, maintenance and functional testing and the related instructions for use.

If class IIa devices are covered by this certificate, the quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis. The devices conform to the technical documentation. The periodical surveillance includes further assessment of the technical documentation on the basis of representative samples.

If class IIb or class III (excluding custom-made implantable devices) devices are covered by this certificate, the quality management system ensures that devices conform to the type that has undergone a type examination. An EU Type-Examination Certificate in accordance with Annex X is required before placing them on the market.

All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with.

For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:G26 057924 0020 Rev. 00

Report No.:

SH2303302

Valid from:

2026-07-28

Valid until:

2031-07-27

Christoph Dicks

Head of Certification/Notified Body

Issue date: 2026-07-28



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Regulation (EU) 2017/745 on Medical Devices, Annex XI Part A

Certificate No. G26 057924 0020 Rev. 00

Classification: Class I
Device Group: H02010106 - METAL SURGICAL STAPLE REMOVERS, SINGLE-USE
Device Properties: MDS 1005 - Devices in sterile condition

Classification: Class IIa
Device Group: MDN 1204 - Non-active non-implantable devices for wound and skin care
Intended Purpose: Disposable skin stapler is intended to be used for skin closure in surgical operations.

The validity of this certificate depends on conditions and/or is limited to the following: N.A.

Revision History:

Rev.	Dated	Report	Description
00	2026-07-28	SH2303302	Initial issuance