



EU Quality Assurance Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex XI Part A
(Class I Devices in sterile condition, with measuring function or reusable surgical instruments)

No. G21 079078 0016 Rev. 00

Manufacturer:

Hangzhou Formed Medical Devices Co., Ltd.

Building 3, No. 292 Gonghe North Road, Xindeng Town
Fuyang District
311404 Hangzhou
PEOPLE'S REPUBLIC OF CHINA

SRN Manufacturer - CN-MF-000011212

Authorized Representative:

Shanghai International Holding Corp. GmbH (Europe)
Eiffestraße 80, 20537 Hamburg, GERMANY

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has established, documented and implemented a quality management system as described in Article 10 (9) of the Regulation (EU) 2017/745 on medical devices. Details on device categories covered by the quality management system are described on the following page(s).

The Report referenced below summarises the result of the assessment and includes reference to relevant CS, harmonized standards and test reports. The conformity assessment has been carried out according to Annex XI Part A of this regulation with a positive result.

As applicable the involvement of the notified body is limited to the 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.

The certified quality assurance system is subject to periodical surveillance by TÜV SÜD Product Service GmbH. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:G21 079078 0016 Rev. 00

Report No.:

SH221041MDR01

Valid from:

2023-07-18

Valid until:

2028-07-17

Christoph Dicks

Head of Certification/Notified Body

Issue date: 2023-07-18



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Classification: Class I
Device Group: R010101 - NASOPHARYNGEAL TUBES
R010102 - GUEDEL AIRWAY TUBES
R018004 - INTUBATION DEVICE GUIDES
Device Properties: MDS 1005.1 - Ethylene Oxide sterilization

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

Revision History:

Rev.	Dated	Report	Description
00	2023-07-18	SH221041MDR01	Initial issuance