



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

Production Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex V

(Devices in Class IIa, IIb or III)

**No. G2 066149 0017 Rev. 01**

## Manufacturer:

**HANGZHOU FUSHAN MEDICAL  
APPLIANCES CO., LTD.**

No. 1288 South Jinxi Road

Linglong Industrial Park

Lin'an District

311301 Hangzhou City, Zhejiang Province

PEOPLE'S REPUBLIC OF CHINA

## Facility(ies):

HANGZHOU FUSHAN MEDICAL APPLIANCES CO., LTD.

No. 1288 South Jinxi Road, Linglong Industrial Park, Lin'an

District, 311301 Hangzhou City, Zhejiang Province, PEOPLE'S

REPUBLIC OF CHINA

## Product

## Category(ies):

**Silicone Urethral Catheter, Trocar, Laryngeal Mask Airway,  
Nebulizer, Stomach Tube, Nasogastric Tube, Drainage Tube,  
Endotracheal Tube, Supra Laryngeal Airway, Mask, Suction  
Tube Kit, Rectal Tube, Oxygen Tube, Yankauer Handle,  
Drainage Kit, Resuscitator, Guedel/Oropharyngeal Airway,  
Intravenous Catheter, Infusion Access Adapter, Gastrostomy  
Tube, Pessary**

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

**Report No.:**

SH1951212

**Valid from:**

2019-10-28

**Valid until:**

2023-07-15

**Date,**

2019-10-28

Stefan Preiß

Head of Certification/Notified Body