

A1 / 04.11



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 16 09 77591 012

Manufacturer:

Hitec Medical Co., Ltd.

No. 1328, Hengnan RD, Minhang District

201114 Shanghai

PEOPLE'S REPUBLIC OF CHINA



EC-Representative:

Shanghai International Holding

Corp. GmbH (Europe)

Eiffestraße 80 20537 Hamburg **GERMANY**

Product Category(ies): Urethral Catheters, Tracheostomy Tube,

Silicone Foley Catheter,

Foley Catheter with Temperature Sensor

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.:

SH16709EXT01

Valid from:

2016-11-03

Valid until:

2021-11-02

Date, 2016-10-10

Stefan Preiß



TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

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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 16 09 77591 012

Facility(ies):

Hitec Medical Co., Ltd.

No. 1328, Hengnan RD, Minhang District, 201114 Shanghai, PEOPLE'S REPUBLIC OF CHINA



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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 16 09 77591 013

Manufacturer:

Hitec Medical Co., Ltd.

No. 1328, Hengnan RD, Minhang District

201114 Shanghai

PEOPLE'S REPUBLIC OF CHINA



EC-Representative:

Shanghai International Holding

Corp. GmbH (Europe)

Eiffestraße 80 20537 Hamburg GERMANY

Product Category(ies):

For detailed information please see

attachment

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.:

SH16709EXT01

Valid from: Valid until: 2016-11-03 2021-11-02

Date. 2016-10-11

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Stefan Preiß



TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123 Page 1 of 3



Attachment for Certificate No G2 16 09 77591 013

Supplement 001 dated 2016-10-11



For the product(s)/product category (ies):

Tracheal Tube, Oxygen Mask, Connecting Tube with Yankauer Handle, Laryngeal Mask Device, Intubating Stylet, Non-rebreath Mask, Tracheostomy Mask, Aerosol Mask, Multi-vent Mask, Stomach Tube, Silicone Stomach Tube, Suction Catheter, Feeding Tube, Nelaton Catheter, Tracheobronchial Tube, Reinforced Endotracheal Tube, Endotracheal Tube Introducer, Nasal Oxygen Cannula, Nebulizer, Disposable Air Cushion Face Mask, Endotracheal Tube Kit, Disposable Breathing Circuits, Heat and Moisture Exchange Filter, Manual Resuscitator, Silicone Tube, Endobronchial Blocker Tube, Ureteral Stent Set, Drainage System, Silicone Drainage System, Endotracheal Tube with Evacuation Lumen, Safety Self-destructive Syringe (with Needle), Sterile Insulin Syringes for Single Use, Disposable Transfusion Set (with Needle), Sterile Hypodermic Syringes for Single Use (with Needle), Disposable Sterile Hypodermic Needles, Infusion Sets for Single Use (with Needle), Scalp Vein Sets

Munich, 2016-10-11

Stefan Preiß

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Product Service

Production Quality Assurance System
Directive 93/42/EEC on Medical Devices (MDD), Annex V
(Devices in Class IIa, IIb or III)

No. G2 17 08 77591 016

Manufacturer:

Hitec Medical Co., Ltd.

No. 703, Hengnan RD 1328

Minhang District 201114 Shanghai

PEOPLE'S REPUBLIC OF CHINA



EC-Representative:

Shanghai International Holding

Corp. GmbH (Europe)

Eiffestraße 80 20537 Hamburg GERMANY

Product Category(ies):

Tracheal Tube, Oxygen Mask,

Connecting Tube (Yankauer Handle) (more details see attachment)

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.:

SH1770907

Valid from: Valid until: 2017-10-23 2021-11-02

Date. 2017-10-23

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ERTIFIKAT ◆ CERTIFICATE

Attachment for Certificate No G2 17 08 77591 016



Product Service

Supplement 001 dated 2017-10-23

For the product(s)/product category (ies):

Laryngeal Mask Device, Intubating Stylet, Non-rebreath Mask, Tracheostomy Mask, Aerosol Mask, Multi-vent Mask, Stomach Tube, Silicone Stomach Tube, Suction Catheter, Feeding Tube, Netaton Catheter, Tracheobronchial Tube, Reinforced Endotracheal Tube, Endotracheal Tube Introducer, Nasal Oxygen Cannula, Nebulizer, Disposable Air Cushion Face Mask, Endotracheal Tube Kit, Disposable Breathing Circuits, Heat and Moisture Exchange Filter, Manual Resuscitator, Silicone Tube, Endobronchial Blocker Tube, Ureteral Stent Set, Drainage System, Silicone Drainage System, Endotracheal Tube with Evacuation Lumen, Safety Self-destructive Syringe (with Needle), Sterile Insulin Syringes for Single Use, Disposable Transfusion Set (with Needle), Sterile Hypodermic Syringes for Single Use (with Needle), Disposable Sterile Hypodermic Needles, Infusion Sets for Single Use (with Needle), Scalp Vein Sets, Sterile Hemodialysis Blood Circuits for Single Use, **Closed Suction System**

Munich, 2017-10-23

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Stefan Preiß

Certification Medical Technology

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EC Certificate

Production Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex V (Devices in class I in sterile conditions, sterilised systems or procedure packs)

No. G2S 17 08 77591 017

Manufacturer:

Hitec Medical Co., Ltd.

No. 703, Hengnan RD 1328

Minhang District 201114 Shanghai

PEOPLE'S REPUBLIC OF CHINA



EC-Representative:

Shanghai International Holding

Corp. GmbH (Europe)

Eiffestraße 80 20537 Hamburg GERMANY

Product Category(ies):

Oropharyngeal Airway, Disposable Rectal Tube, Nasopharyngeal Airway, Urine Bag, Spigot

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for manufacture in accordance with MDD Annex V. This quality assurance system covers those aspects of manufacture concerned with securing and maintaining sterile conditions of the respective devices / device categories and conforms to the requirements of this Directive. It is subject to periodical surveillance. See also notes overleaf.

Report No.:

SH1770907

Valid from: Valid until: 2017-10-23 2021-11-02

Date, 2017-10-23

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TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123 Page 1 of 2

TÜV SÜD Product Service GmbH · Zertifizierstelle · Ridlerstraße 65 · 80339 München · Germany





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 17 08 77591 018

Manufacturer:

Hitec Medical Co., Ltd.

No. 703, Hengnan RD 1328

Minhang District 201114 Shanghai

PEOPLE'S REPUBLIC OF CHINA



EC-Representative:

Shanghai International Holding

Corp. GmbH (Europe)

Eiffestraße 80 20537 Hamburg **GERMANY**

Product

Urethral Catheters, Tracheostomy Tube,

Silicone Foley Catheter, Category(ies):

Foley Catheter with Temperature Sensor

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 .:

SH1770907

Valid from:

2017-10-23

Valid until:

2021-11-02

Date, 2017-10-23

Stefan Preiß

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