



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 065758 0004 Rev. 01

Manufacturer:

Shenzhen Biocare Bio-Medical

Equipment Co., Ltd.

#16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District

Pingshan New District 518122 Shenzhen

PEOPLE'S REPUBLIC OF CHINA

Facility(ies):

Shenzhen Biocare Bio-Medical Equipment Co., Ltd. #16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District,

Pingshan New District, 518122 Shenzhen, PEOPLE'S REPUBLIC

OF CHINA

Product Category(ies): Digital Electrocardiograph, Patient Monitor, **B-Ultrasonic Diagnostic Equipment,**

Doppler Fetal Heart Rate Detector, Infusion

Pump, Syringe Pump, Fingertip Pulse Oximeter, Handheld Pulse Oximeter,

Fetal/Maternal Monitor, Fetal Monitor, Color

Doppler Ultrasound System, Central Monitoring System, Ambulatory

Electrocardiographs, Ambulatory blood

pressure recorders, and associated

software.

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.:

BJ1989607

Valid from:

2019-09-11

Valid until:

2024-05-26

Date,

2019-09-11

Stefan Preiß

1. Punil

Head of Certification/Notified Body

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