

EC Certificate

Directive 93/42/EEC Annex II, excluding Section 4 Full Quality Assurance System Medical Devices

Registration No.: HD 60144433 0001

Report No.:

17054024 008

Manufacturer:

Shenzhen Upnmed Equipment

Co., Ltd.

4th Floor, Building #1 East **Huihuang Industrial Area**

Xitian Community, Gongming Town

Guangming New District

Shenzhen

518107 Guangdong

Products:

China

Oximeter Probes

Replaces Approval, Registration No.: HD 60126823 0001

Expiry Date:

2024-05-27

The Notified Body hereby declares that the requirements of Annex II, excluding section 4 of the directive 93/42/EEC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex II, section 5 of the aforementioned directive. For placing on the market of class III devices covered by this certificate an EC design-examination certificate according to Annex II, section 4 is required.

Effective Date:

2019-12-02

Date:

2019-12-02

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 93/42/EEC concerning medical devices with the identification number 0197.