

| Manufacturer                                                                                                                                            | Authorized European Representative                                               | Notified Body                                                                          |
|---------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|
| Covidien llc<br>15 Hampshire Street<br>Mansfield, MA 02048 USA<br>(formerly: Nellcor Puritan Bennett L.L.C., a<br>division of Tyco Healthcare Group LP) | Covidien Ireland Limited<br>IDA Business & Technology Park<br>Tullamore, Ireland | TÜV SÜD Product Services GmbH<br>Ridlerstrasse 65<br>D-80339 Munich<br>Germany<br>0123 |

## Declaration of Conformity

**Document #/Revision #:** 10138709, Rev. N

**Product/Family Name:** Nellcor Bedside SpO<sub>2</sub> Patient Monitoring System (PM100N)

**Classification Rationale:** Class IIb per Rule II of Annex IX and Class 1 per Rule 1 of Annex IX

**EU Conformity Assessment Route:** Class IIb products according to Annex II, Class I products according to Annex VII

**Standards Applied:** A complete listing of the applied standards is available in Section 4 of the Nellcor Bedside SpO<sub>2</sub> Patient Monitoring System (Big Bird) Technical File Part A p/n 10138705

**Start of CE Marking:** 01/ 2015

Covidien llc declares under our sole responsibility that the above product(s) to which this declaration relates, and which bear(s) the CE Marking, is (are) in conformity with the Essential Requirements of EC Directive 93/42/EEC of 14 June 1993, as amended by 2007/47/EC of the European Parliament and of the Council, concerning medical devices, which allows their free distribution, sale and circulation in the European Union (EU); they comply with the provisions of the defined regulatory requirements and which comply with the referenced standards, as stated above.

This declaration is made in accordance with the requirements of Clause 1.8 of Schedule 3 of the Australian Therapeutic Goods (TGA) Medical Device Regulations 2002, relating to the devices stated in Schedule I of this document.

Covidien llc hereby declares that all medical devices referenced in Schedule I placed on the European Community market by the Company & its subsidiaries are compliant with Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the Restriction of Certain Hazardous Substances in Electrical and Electronic Equipment (commonly known as the EU RoHS Directive). They are RoHS compliant.

- All supporting documentation is retained by the manufacturer
- As required by the above Directive, this Declaration is supported by
  - EC Certificate: MDD Annex II, G1 077790 0060 Rev 00, Reference P/N 10069299, issued by TÜV SÜD Product Services GmbH, Ridlerstrasse 65, D-80339 Munich Germany, on 29 June 2020
- This Declaration of Conformity is applicable to all of the medical devices referenced in Schedule I, manufactured by Covidien llc and/or produced under its certified Quality System control. Products referenced in Schedule I can be traced by means of the related product identification referenced in the relevant labeling (i.e.: lot number, serial number, etc.).
- Each kind of medical device to which the Full Quality Assurance Procedures have been applied complies with the applicable provisions of the essential requirements/principles, the classification rules, at each stage, from the design of the device until its final inspection before being supplied.

This Declaration shall be retained for a period of the lifetime of the medical device (LMD) + 1 year or minimum of 15 years once the record is obsoleted or superseded.

**Date of Issue:** 16 Mar 2021

**Place of Issue:** Boulder, Colorado, USA

**Signature:**

*Greeshma Kayala*

Signer Name: Greeshma Kayala  
Signing Reason: I approve this document  
Signing Time: 16 March 2021 | 13:54 CDT

56620C7713D74F81BFBF217451CDD1E3

**Name/Title** Greeshma Kayala, Regulatory Affairs Manager

**Schedule 1**  
**Declaration of Conformity for**  
**Nellcor Bedside SpO<sub>2</sub> Patient Monitoring System (Big Bird)**

| Medical Device Part Number | Description                                                | Class/Rule | UMDNS Code and Term      | GMDN Code and Term |
|----------------------------|------------------------------------------------------------|------------|--------------------------|--------------------|
| PM100N                     | Nellcor Bedside SpO <sub>2</sub> Patient Monitoring System | IIb/10     | 17148 – Oximeters, Pulse | 17148 - Oximeters  |

| Accessory Part Number | Description                                   | Class/Rule | UMDNS Code and Term                               | GMDN Code and Term                                |
|-----------------------|-----------------------------------------------|------------|---------------------------------------------------|---------------------------------------------------|
| DOC10                 | Nellcor Pulse Oximetry Interface Cable DOC-10 | I          | 17603 – Extension cords, electrical, multi-outlet | 36092 – Cable, electrical extension, multi-outlet |