
Declaration of Conformity

We declare under our sole responsibility that the following product to which this declaration relates is in conformity with the standards or other normative documents under the provisions of Medical Devices Regulation (EU) 2017/745 and RoHS Directive 2011/65/EU.

Product Category: Measuring and Monitoring Device for Vital Signs
Device Name: DYNASCOPE 7000 series Central Monitor DS-7700 system
(Refer to the Appendix 1 for the details of the system components)
Classification: Class IIb
Rule 10, Annex VIII of Medical Devices Regulation (EU) 2017/745
Basic UDI-DI: 4538612021009S
Intended Purpose: This device is a central monitor to monitor the conditions of the patients. It deals with the following vital sign parameters, which are measured and transmitted by the specified bedside monitors and /or the telemeters for use in combination.
The vital signs are electrocardiogram, heart rate, respiration rate, body temperature, arterial oxygen saturation (SpO₂), pulse rate, invasive blood pressure, non-invasive blood pressure, CO₂ concentration, O₂ concentration and anesthetic gas concentration (including N₂O, halothane, isoflurane, enflurane, sevoflurane and desflurane).
EEE Categories: 8
Annex I of RoHS Directive 2011/65/EU
Manufacturer: Fukuda Denshi Co., Ltd.
3-39-4 Hongo, Bunkyo-ku, Tokyo, Japan
SRN: JP-MF-000008253
European Authorized Representative: Emergo Europe
Prinsessegracht 20, 2514 AP The Hague, The Netherlands
SRN: NL-AR-000000116
Applied Annex: Annex IX Chapter I and III of Medical Devices Regulation (EU) 2017/745
Notified Body: BSI Group The Netherlands B.V. (2797)
Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands
Certificate No.: MDR 743406
Note: BSI's CE Certificate covers only Medical Devices Regulation (EU) 2017/745 Class IIa and IIb, but not RoHS Directive 2011/65/EU.
Technical Documents: No. 063020H-22215

We declare under our sole responsibility that the following product to which this declaration relates is in conformity with the standards or other normative documents under the provisions of RE Directive 2014/53/EU.

Product Category: Measuring and Monitoring Device for Vital Signs
Device Name: DYNASCOPE 7000 series Central Monitor DS-7700 system
Model: DS-7700
Type: DS-7780W, DS-7780, LW-7080
Manufacturer: Fukuda Denshi Co., Ltd.
3-39-4 Hongo, Bunkyo-ku, Tokyo, Japan
Location of design: Fukuda Denshi Co., Ltd. Hongo Office
2-35-8 Hongo, Bunkyo-ku, Tokyo, Japan
Location of production: Fukuda Denshi Co., Ltd. Shiroy Factory
305-1 Naka, Shiroy-shi, Chiba, Japan
Applied Annex: Annex II of Directive 2014/53/EU
Technical file: No. 063020D-17605
No. 063020D-17609

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Quality Assurance Department

Appendix 1 (Details of the system components)

Device Name	Model	Type
DYNASCOPE 7000 series Central Monitor DS-7700 system	DS-7700	DS-7780W
DYNASCOPE 7000 series Central Monitor DS-7700 system	DS-7700	DS-7700WL
DYNASCOPE 7000 series Central Monitor DS-7700 system	DS-7700	DS-7780
DYNASCOPE 7000 series Central Monitor DS-7700 system	DS-7700	DS-7700L
DYNASCOPE 7000 series Central Monitor DS-7700 system	DS-7700	LW-7080