

Declaration of Conformity

The undersigned company

ULTRA VIOL sp. j. Pietras, Purgał, Wójcik 34 Stępowizna Street 95-100 Zgierz POLAND

declares that the product

Product name:

Flow germicidal lamps

NBVE 60; NBVE 60/30; NBVE 110; NBVE 110/55

Mode of suspension:

N-wall mounted; S-ceiling mounted; P-on mobile stand

Conforms the essential requirements stated in the following EC – Directives:

- MDD Directive 93/42/EEC and 2007/47/EC
- EMC Directive 2004/108/EC

Products are medical devices of class I, rule 12 according to Annex IX of Medical Device Directive MDD 93/42/EEC and 2007/47/EC and conforms the essential requirements stated in Annex I of this directive.

The conformity assessment was carried out according to Annex VII of Council Directive 93/42/EEC and 2007/47/EC.

The devices conforms the harmonized European standards:

- EN 60601-1:2006 + A1:2013 Medical electrical equipment Part 1: General requirements for basic safety and essential performance,
- EN 60601-1-2 Medical electrical equipment Part 1-2: General requirements for basic safety and essential performance Collateral standard: Electromagnetic compatibility Requirements and testsand

We declare with full responsibility that the products meet the requirements of the RoHS directive 2011/65/EU (including all its changes and amendments). Conformity assessment was carried out according to standard EN 50581: 2012.

Quality Management System of ULTRA-VIOL certified by TUV Nord meets requirements of:

- EN ISO 9001 Quality Management Principles
- EN ISO 13485 Medical devices Quality management systems.

Wiesław Pietras GENERAL MANAGER

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20.10.2014 r.