

EC DECLARATION OF CONFORMITY

According to annexe VII of the council directive 93/42/EEC regarding medical devices

The manufacturer,






IDMED,

Hôtel Technoptic, 2 rue Marc Donadille
13013 – Marseille – France

Declares and certifies, under its sole responsibility, that the

accessory of the NeuroLight & AlgiScan devices:


with the following references are compliant with:

- the essential requirements of the Directive 93/42/EEC, its amendments and the French Public Health Code
- the following harmonized standards:
 -  IEC 60601-1: 2012: Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
 -  IEC 60601-1-2: 2014: Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance – Collateral Standard: Electromagnetic disturbances – Requirements and tests
 -  IEC 60601-1-6: 2013 - Ed. 3.1 Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability
 -  ISO 14971: 2019 Medical devices - Application of risk management to medical devices
 -  IEC 62366-1:2015-Ed. 1.1 Medical Devices - Part 1: Application Of Usability Engineering To Medical Devices

REFERENCE	DESCRIPTION
STA-W2	Wireless charging station

According to the annex IX of the European directive 93/42/EEC and its amendments, these accessories are Class I rule 12.

The declaration is based on following elements:

-  Technical file DT_NeuroLight & DT_AlgiScan attesting the compliance to the essential requirements of the directive 93/42/EEC

Marseille, 03 November 2020




Frederic BERNERT – President