

## **EU DECLARATION OF CONFORMTIY**

## I.A.C.E.R. S.r.l

Via Enzo Ferrari 2 – 30037 Scorzè (Ve), Italia SRN (Single Registration Number): IT-MF-000009126

herewith declares under its own responsibility, that the product

### **MAG3000**

UDI-DI: 08019781401083 Basic UDI-DI: 8019781PEMFLFDEVP2 Batch no.:

Series no.:

is a magnetotherapy device studied and indicated for the rehabilitation and functional recovery treatments of pathologies affecting:

- wrist, hand, shoulder, foot, ankle and knee joints;
- skeletal motor system;
- muscular atrophies and dystrophies;
- contusions;
- distortions;
- benign injuries and muscle tears;

and for the care treatments of:

- osteoporosis;
- bone edema;
- osteonecrosis;
- ulcers;
- neuropathies;
- arthrosis;
- bunions;
- periarthritis;
- tendonitis and tendinosis.

It complies with the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 5 April 2017 relating to medical devices, which amends directive 2001/83/EC and repeals directives 90/385/ EEC and 93/42/EEC of the Council, and further modifications.





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The device is classified as class IIa, in accordance with regulation 9 (6.1) of Annex VIII to this Regulation and meet all applicable standards and the general safety and performance requirements of the Regulation (EU) 2017/745 Annex 1, and bear the mark



Compliance of the concerned product with the Regulation (EU) 2017/745 has been assessed and certified by the Notified Body

#### 1936 – TÜV Rheiland Italia Via Enrico Mattei 3, 20010 Polignano Milanese (MI), Italia Num. Certificato: ITH 1344294 1

following the certification procedure according to the Regulation (EU) 2017/745, Annex IX.

The devices comply with the following applicable standards:

EN 60601-1:2006/A2:2021, EN 60601-1-2:2015/A1:2021, EN 60601-1-11:2015, EN ISO 60601-1-6: 2010+A1:2015, EN ISO 14971: 2019/A11:2021, EN ISO 10993-1: 2020, EN ISO 10993-5: 2009, EN ISO 10993-10: 2013, EN ISO 10993-18:2020, EN ISO 15223-1: 2021, EN ISO 20417:2021, EN 62304:2006/AC:2008, EN ISO 62366-1:2015+AMD1:2020, ISTA 2A.

It is also claimed that:

- the devices do not incorporate as an integral part any substance or derivative of human blood referred to in point 8 of article 1 of this regulation;
- no fabrics of animal origin have been used in the production as per Regulation (EU) no. 722/2012 of the Commission;
- no common specifications other than the standards mentioned above apply.

It also declares the conformity of the aforementioned products by issuing this EU Declaration of Conformity after having drawn up the technical documentation referred to in Annexes II and III of Regulation (EU) 2017/745 pursuant to article 52 (7) of the Regulation (EU) 2017/745.

Scorzè, 16/01/2023

Place, date

MASSIMO MARCON

Legal Representative

I.A.C.E.R. Srl

#### MD117-08 Data.Rev.31/01/22

Via Enzo Ferrari 2 - 30037 Scorzè (VE) – Italia/Italy - Tel.: (+39) 041/5401356 - Email: iacer@iacer.it PEC: iacer@pec.it - Web: www.itechmedicaldivision.com - Cod. Fisc./P.IVA/VAT N.: ITo0185480274 R.E.A.: VE N. 120250 - M. VE001767 - Codice SDI/SDI Code: SUBM70N - Cap. Soc.: € 1.000.000,00 i.v.

