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## EC DECLARATION OF CONFORMITY

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

Via Enzo Ferrari 2 – 30037 Scorzè (Ve), Italia

herewith declares under its own responsibility, that the product

**LaMagneto X**

UMDNS Code: **12415**

Batch no.:

Series no.:

has been designed and manufactured according to the European Medical Device Directive 93/42/EEC (transposed in Italy by the D.Lgs. 46/97), as modified by the Directive 2007/47/EC (D.Lgs.37/2010) and further modifications/integrations.

The products have been assigned to class IIa, according to Annex IX, rule 9 of the Directive 93/42/EEC (and further modifications/integrations) and bear the mark



Compliance of the concerned products with the Directive 93/42/EEC has been assessed and certified by the Notified Body

**1936 – TÜV Rheinland Italia**

**Via Enrico Mattei 3, 20010 Polignano Milanese (MI), Italia**

Certificate no.: HD60134521

following the certification procedure according to Annex II (excluding point 4) of the Directive 93/42/EEC.

Scorzè, 04/08/2022

*Place, date*



MASSIMO MARCON

*Legal Representative*

**MD117-07 Date.Rev.31/01/22**

**I.A.C.E.R. Srl**

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PEC: iacer@pec.it - Web: www.itechmedicaldivision.com - Cod. Fisc./P.IVA/VAT N.: IT00185480274  
R.E.A.: VE N. 120250 - M. VE001767 - Codice SDI/SDI Code: SUBM70N - Cap. Soc.: € 1.000.000,00 i.v.

