

## EC DECLARATION OF CONFORMTIY

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

Via S.Pertini 24/A - 30030 Martellago (Ve), Italia

herewith declares under its own responsibility, that the product

**MIO CARE PRO** 

**UMDNS Code: 13762** 

Batch no.:

Series no.:

comply with the essential safety requirements of 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 product has been assigned to class IIa, according to Annex IX, rule 9 of the Directive 93/42/EEC (and further modifications/integrations) and bears the mark



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

0068 – MTIC InterCert S.r.l.

Via G. Leopardi 14, Milano (MI) 20123, Italia
Certificate no.: 0068/QCO-DM/234-2020

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

The devices comply with the following applicable standards:

EN 60601-1:2006 + A1:2013, EN 60601-1-2:2015, IEC 60601-1-6:2013, EN 60601-1-11:2015, IEC 60601-2-10:2012, EN ISO 14971: 2012, ISO 10993-1: 2009+AC:2010, ISO 10993-5: 2009, ISO 10993-10: 2010, IEC 62133:2012, IEC 62304:2015, EN 62366-1:2015.

It is also claimed that:

- The devices do not incorporate, as an integral part, any substance or a human blood derivate of point 10 of Annex 1 (Directive 2007/47/CE);
- tissue of animal origin, provided for in Directive 2003/32/CE, haven't been used in production.

Martellago, 03/08/2020

Place, date

MASSIMO MARCON

Legal Representative