

Declaration of Conformity

For the following products:

Electrical Stimulator

(Product Name)

R-C1, R-E1, R-T1

(Model Designation)

is hereinafter confirmed to comply with the requirements set out in the Council Directive on the harmonization of the Laws of the Member States concerning Medical Device Directive (93/42/EEC As amended by 2007/47/EC)

EN ISO 13485:2016

EN ISO 15223-1:2016

EN 1041:2008

EN ISO 780:2015

EN ISO 10993-1:2009/AC:2010

EN ISO 10993-5:2009

EN ISO 10993-10:2010

EN ISO 14971:2012

EN 60601-1:2006/A1:2013

EN 60601-1-2:2015

EN 60601-1-6:2010+A1:2015

EN 60601-1-11:2015

EN 60601-2-10:2015+A1:2016

EN 62304:2006/A1:2015

EN 62366-1:2015

Conformity Assessment Route:

Annex II excluding section 4 of Medical Device Directive

Notified Body:

DNV GL Presafe AS (NB No. 2460)

Veritasveien 3, N-1363 Høvik Postbox 116, N-1300 Sandvika

The following representative in Europe is responsible for making this declaration:

Company Name: Shanghai International Holding Corp. GmbH (Europe)

Company Address: Eiffestrasse 80, 20537 Hamburg, Germany

The following manufacturer is responsible for making this declaration:

Company Name: Shenzhen Roundwhale Technology Co., Ltd.

Company Address: 202, 2/F, Building 27, Dafa Industrial Park, Longxi Community, Longgang District, Shenzhen 518000, China.



Vice President

(Position/title)

2020.6.16

(Date)