

Declaration of Conformity

Product Description	Product Name	Class
Pulse Oximeter	Unit: SA110, SA120, SA200, SA210, SB100, SB200, SB210, SB220, SA300, SA310, SA320, SD100 Probe: PA100, PB100, PC100, PD100, PF100	IIB

Is herewith confirmed to comply with the requirements set out in the Council Directive on the harmonization of the laws of Member States concerning **Medical Devices Directive 93/42/EEC as amended by 2007/47/EC** with the compliance of conformity assessment Annex II-exclusive section 4 to be certified by **DNV GL Presafe AS (notify body number –2460)**.

For the evaluation regarding the **Class IIB** product safety aspects, the following harmonized standards are applied: EN ISO13485:2016; ISO80601-2-61:2011; EN ISO14971:2012; EN ISO15223-1:2016; EN 1041:2008; EN 60601-1:2006/A1:2013; EN 60601-1-2:2015; EN 60601-1-6:2010 ; EN 60601-1-11:2010; EN ISO10993-1:2009; EN ISO10993-5:2009 ; ISO10993-10:2010 ; EN 62366:2008; EN 62304:2006

The following European Authorized Representative is stated to the declaration:

Representative Name: **CMC Medical Devices & Drugs S.L.**

Representative Address : **C/ Horacio Lengo Nº 18, CP 29006, Málaga, Spain**

The following person is responsible for the compliance of declaration:

Manufacturer Name : **Rossmax Innotek Corp**

Head office : **12F., No. 189, Kang Chien Rd, Taipei 114, Taiwan**

Manufacturing site: **1F/6F., No. 789, Bo-Ai St., Jhubei City, Hsinchu County 302, Taiwan.**

President

(Position / Title)



(Legal Signature)

May 22, 2019

(Date)