File Name: Declaration of Conformity File No.: CS/CE-MD300CN310-01

ChoiceMMed Revision: E

Page 1 of 3

Declaration of Conformity
to Council Directive 93/42/EEC
concerning Medical Devices

concerning medical Devices	
Manufacturer:	Beijing Choice Electronic Technology Co., Ltd.
	Room 4104, No. A12 Yuquan Road, Haidian District, 100143 Beijing, PEOPLE'S REPUBILIC OF CHINA.
European Representative:	Shanghai International Holding Corp. GmbH (Europe)
	Eiffestraße 80 20537 Hamburg GERMANY
Product Name:	Fingertip Pulse Oximeter
Product Model:	See attached list
UMDNS Code:	17148
Classification:	Class IIa, rule 10 to Annex IX of the MDD
Conformity assessment Route:	Annex II excluding (4)
We, the manufacture	, herewith declare that the stated medical devices
meet the transposition in	nto national law, the provisions of Council Directive
93/42/	EEC concerning medical devices.
All supporting document	ation is retained at the premises of the manufacturer.
Standards applied:	
EN ISO 13485:2016/AC:2016 N	Iedical devices- Quality management systems-
Requirements for regulatory pur	poses
EN ISO14971:2012 Medical dev	vices - Application of risk management to medical devices
EN 60601-1:2006/A1:2013 Med	ical electrical equipment-Part 1: General requirements for
safety	
EN 60601-1-2:2015 Medical electrical equipment Part 1-2: General requirements for basic	
safety and essential performance - Collateral Standard: Electromagnetic disturbances -	
Requirements and tests	
EN 60601-1-6:2010 Medical ele	ctrical equipment Part 1-6: General requirements for
basic safety and essential perform	nance - Collateral standard: Usability
EN 60601-1-11:2010 Medical el	ectrical equipment Part 1-11: General requirements for



basic safety and essential performance - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment. ISO 80601-2-61:2011 Medical electrical equipment —Part 2-61: Particular requirements for basic safety and essential performance of pulse oximeter equipment EN ISO10993-1:2009/AC:2010 Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process EN ISO10993-5:2009 Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity ISO10993-10:2010 Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization EN1041:2008 Information supplied by the manufacture of medical devices EN ISO15223-1:2016 Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements EN 62304:2006/AC:2008 Medical device software-Software life-cycle processes MEDDEV 2.7/1: 2016 CLINICAL EVALUATION: A GUIDE FOR MANUFACTURERS AND NOTIFIED BODIES UNDER DIRECTIVES 93/42/EEC and 90/385/EEC TÜV SÜD Product service GmbH Notified Body: Ridlerstr 65, D-80339 München, Germany **CE** 0123 Identification Number: (EC) Certificate(s): No. G1 078179 0032 Rev.01 Start of CE-marking: 2016-05-06

Place, Date of Declaration:

Signature:

Sile

Name: Haiying Zhao

Beijing, 2019-05-22

Position: Quality Director



Attached list

MD300CN310, MD300CN330, MD300CN340, MD300CN350, MD300CN356, MD300CN360, MD300CN130, MD300CN150, MD300CN160

MD300C1, MD300C11, MD300C12, MD300C13, MD300C15, MD300C16, MD300C17, MD300C18, MD300C19, MD300C1B, MD300C1C, MD300C1D, MD300C1E, MD300C1F, MD300C15D

MD300C2, MD300C20, MD300C201, MD300C203, MD300C204, MD300C21, MD300C21C, MD300C22, MD300C221, MD300C23, MD300C25, MD300C26, MD300C29, MD300C2A, MD300C2B, MD300C2D, MD300C2E, MD300C2F

MD300C4, MD300C41

MD300C5, MD300C52, MD300C53, MD300C54

MD300C63, MD300C634, MD300CF3, MD300CH3

LTD800, LTD805