**ChoiceMMed** File Name: Declaration of Conformity File No.: CS/CE-MD300C29-H-01

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Decla	aration of Confo	rmity
to Cour	ncil Directive 93/	<b>/42/EEC</b>
conce	rning Medical D	Devices
Manufacturer:	Beijing Choice Electronic Technology Co., Ltd.	
	Room 4104, No.A12 Yuquan Road, Haidian District,	
	100143 Beijing, P.R.CHINA	
European Representative:	Shanghai International Holding Corp. GmbH (Europe) Eiffestraβe 80 20537 Hamburg GERMANY	
Product:	Fingertip Pulse Oximeter	
	Customers type no.:	Manufactures type no.:
	Microlife OXY300	MD300C2G/MD300C29
UMDNS Code:		17148
Classification:	Class IIa, rule 10 to Annex IX of the MDD	
Conformity assessment Route:	Annex 1	I excluding (4)
meet the transposition 93/4	er, herewith declare that the s into national law, the provisi 2/EEC concerning medical de ntation is retained at the prem	ons of Council Directive evices.
meet the transposition 93/4 All supporting documen	into national law, the provisi 2/EEC concerning medical de	ons of Council Directive evices.
meet the transposition 93/4 All supporting documen Standards applied:	into national law, the provisi 2/EEC concerning medical dentation is retained at the prem	ons of Council Directive evices. lises of the manufacturer.
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical d	into national law, the provisi 2/EEC concerning medical dentation is retained at the prem	ons of Council Directive evices. lises of the manufacturer.
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical d regulatory purposes	into national law, the provisi 2/EEC concerning medical dentation is retained at the preme evices- Quality manageme	ons of Council Directive evices. lises of the manufacturer. nt systems- Requirements for
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical d regulatory purposes EN ISO14971:2012 Medical de	into national law, the provisi 2/EEC concerning medical dentation is retained at the preme evices- Quality manageme	ons of Council Directive evices. lises of the manufacturer. nt systems- Requirements for
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical d regulatory purposes EN ISO14971:2012 Medical de devices	into national law, the provisi 2/EEC concerning medical dentation is retained at the preme evices- Quality manageme evices – Application of risk	ons of Council Directive evices. dises of the manufacturer. nt systems- Requirements for management to medical
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical d regulatory purposes EN ISO14971:2012 Medical de devices EN ISO10993-1: 2009/AC:201	into national law, the provisi 2/EEC concerning medical dentation is retained at the preme evices- Quality manageme evices – Application of risk 0 Biological evaluation of	ons of Council Directive evices. dises of the manufacturer. nt systems- Requirements for management to medical
meet the transposition 93/4 All supporting documen Standards applied: EN ISO 13485:2012 Medical do regulatory purposes EN ISO14971:2012 Medical do devices EN ISO10993-1: 2009/AC:201 Evaluation and testing within a	into national law, the provisi 2/EEC concerning medical dentation is retained at the preme evices- Quality manageme evices – Application of risk 0 Biological evaluation of risk management system	ons of Council Directive evices. dises of the manufacturer. nt systems- Requirements for management to medical

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File Name: Declaration of Conformity File No.: CS/CE-MD300C29-H-01

EN ISO10993-10:2010 Biological Evaluation of Medical Devices-Part 10: Tests for irritation and delayed-type hypersensitivity

EN 60601-1:2006/A1:2013 Medical electrical equipment-Part1: General requirements for safety and essential performance

EN 60601-1-2:2007/AC:2010 Medical electrical equipment-Part1-2: General

requirements for safety and essential performance Collateral Standard: Electromagnetic compatibility – Requirements and tests

ISO 80601-2-61:2011 Medical electrical equipment-Particular requirements for basic

safety and essential performance of pulse oximeter equipment for medical use

EN1041:2008 Information supplied by the manufacture of medical device

EN 980:2008 Symbols for use in the labelling of medical devices

EN 62304: 2006 Medical device software-Software life-cycle processes

EN60601-1-6: 2010 Medical electrical equipment-Part 1-6: General requirements for basic safety and essential performance-Collateral Standard: Usability

Notified Body:

TÜV SÜD Product service GmbH Ridlerstr 65, D-80339 München, Germany



Identification Number:

(EC) Certificate(s):

Start of CE-marking:

Place, Date of Declaration:

No. G1 16 02 78179 025

2014-02-14

Beijing, 2017-01-13

Lez Chen

Name: Lei Chen Position: Quality Director

Signature: