## **EC DECLARATION OF CONFORMITY**

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## **EC DECLARATION OF CONFORMITY**

Name and address of the manufacturer:	GuangZhou AXD Electronic Co., Ltd. Factory 3, No.9 Changxin Road, Changshapu, Zhongluotan Town, 510550, Baiyun District, Guangzhou City, Guangdong Province, P.R.China
Name and address of the European Representative	Shanghai International Holding Corp. GmbH (Europe) Eiffestrasse 80, 20537 Hamburg, Germany
We declare under our sole responsibility that	
the medical device:	<u>Digital Arm Blood Pressure Monitor</u> including (Product Name)
	AXD-806, AXD-808, AXD-809 (Model Designation)
of class:	IIa according to rule 10 in annex IX of directive 93/42/EEC
meets the provisions of the directive 93/42/EE	EC and its transpositions in national laws which apply to it.
Conformity assessment procedure:	Directive 93/42/EEC Annex V
Registration No.:	FI21/07010
Notified Body:	SGS Fimko Ltd. Takomotie 8, FI-00380 Helsinki, Finland CE0598
	Moore
Guangzhou, May 21, 2021	General Manager
Place, date	Name and function

Applied Standards are listed as below:

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No.	Standard No.	Standard Description	
1	EN ISO 13485:2016	Medical devices – Quality management systems - Requirements for regulatory purposes	
2	EN 60601-1:2006+A1:2013	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance	
3	EN 60601-1-2:2015	Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests	
4	EN 80601-2-30:2010+A1:2015	Medical electrical equipment – Part 2-30: Particular requirements for the basic safety and essential performance of automated non-invasive sphygmomanometers	
5	EN 1060-1:1995+A2:2009	Non-invasive sphygmomanometers – Part 1: General requirements	
6	EN 1060-3:1997+A2:2009	Non-invasive sphygmomanometers – Part 3: Supplementary requirements for electro-mechanical blood pressure measuring systems	
7	EN ISO 81060-2:2019	Non-invasive sphygmomanometers - Part 2: Clinical validation of automated measurement type	
8	EN 60601-1-11:2015	Medical electrical 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	
9	EN ISO 14971:2012	Medical devices – Application of risk management to medical devices	
10	EN ISO 10993-1:2009+AC:2010	Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process	
11	EN ISO 10993-5:2009	Biological evaluation of medical devices – Part 5: Tests for in vitro cytotoxicity	
12	EN ISO 10993-10:2013	Biological evaluation of medical devices – Part 10: Tests for irritation and skin sensitization	
13	EN 62304:2006+A1:2015	Medical device software - Software life-cycle processes	
14	EN 60601-1-6:2010+A1:2015	Medical electrical equipment Part 1-6: General requirements for basic safety and essential performance - Collatera standard: Usability	
15	EN 62366-1:2015	Medical devices – Part 1: Application of usability engineering to medical devices	
16	EN 1041:2008+A1: 2013	Information supplied by the manufacturer of medical devices	
17	EN ISO 15223-1:2016	Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements	
18	ISTA 2A:2011	Performance Test for Packaged-Products weighing 150 lb (68 kg) or Less	