DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 93/42/EEC CONCERNING MEDICAL DEVICES

CONTEC MEDICAL SYSTEMS CO., LTD No.112 Qinhuang West Street, Economic & Technical MANUFACTURER: Development Zone, Qinhuangdao, Hebei Province, PEOPLE'S REPUBLIC OF CHINA MEDICAL DEVICE: SPIROMETER, SP10 **CLASSIFICATION - ANNEX IX:** Class II a, Rule 10 CONFORMITY ASSESSMENT ROUTE: Annex II excluding chapter 4 WE, (CONTEC MEDICAL SYSTEMS CO., LTD) HEREWITH DECLARE THAT THE STATED DEVICES. MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE 93/42/EEC of 14 JUNE 1993 CONCERNING MEDICAL DEVICES; INCLUDING, AT 21 MARCH 2010, THE AMENDMENTS BY COUNCIL DIRECTIVE 2007/47/EEC ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURE. STANDARDS APPLIED: SEE ATTACHED LIST OF (HARMONISED - EN) STANDARDS FOR WHICH DOCUMENTED EVIDENCE OF COMPLIANCE CAN BE PROVIDED. TÜV SÜD PRODUCT SERVICE GMBH NOTIFIED BODY: RIDLERSTR 65, D-80339 M NCHEN, GERMANY **C** € ₀₁₂₃ **IDENTIFICATION NUMBER:** (EC) CERTIFICATE(S): G1 050972 0050 Rev.02 EC REP Shanghai International Holding Corp. GmbH(Hamburg) **EUROPEAN REPRESENTATIVE:** Eiffestrasse 80, 20537 Hamburg Germany START OF CE-MARKING: 2011-01-14 (Date or Lot or serial number) PLACE, DATE OF DECLARATION: QINHUANGDAO, 2019-07-23 SIGNATURE:

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Appendix: list of (harmonised - EN) standards

NO.	Reference	Title
1	EN60601-1:1990+A1:1993+A2:1995	Medical electrical equipment- Part 1:
		General requirements for safety
2	EN 60601-1-2:2007	Medical electrical equipment- Part 1-2:
		General requirements for safety -
		Collateral standard: Electromagnetic
		compatibility - Requirements and tests
3	EN60601-1-4:1996+A1:1999	Medical electrical equipment - Part 1-4:
		General requirements for safety -
		Collateral standard: Programmable
		electrical medical systems
4	EN 60601-1-6:2007	Medical electrical equipment-Part 1-6:
		General requirements for basic safety and
		essential performance-Collateral
		Standard: Usability
5		Anaesthetic and respiratory equipment -
	EN ISO23747:2009	Peak expiratory flow meters for the
	(ISO 23747:2007)	assessment of pulmonary function in
		spontaneously breathing humans
6	EN 62304:2006	Medical device software-Software
		life-cycle processes

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