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

Meditech Equipment Co.,Ltd

89 Laoshan Road, Building 69, MANUFACTURER: Laoshan District, Qingdao, Shandong Province, China **MEDICAL DEVICE:** MD908B Patient Monitor. **CLASSIFICATION - ANNEX IX:** Class Ilb. Rule 10 **CONFORMITY ASSESSMENT ROUTE:** Annex IX excluding chapter 4 WE, (MEDITECH EQUIPMENT CO., LTD) HEREWITH DECLARE THAT THE STATED MEDICAL 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. NOTIFIED BODY: MEDCERT GmbH Pilatuspool 2. 20355 Hamburg, GERMANY **IDENTIFICATION NUMBER:** (EC) CERTIFICATE(S): 7368GB410191111 OBELIS S. A **EUROPEAN REPRESENTATIVE:** Registered Address: Bd. Général Wahis, 53 1030 Brussels, Belgium Tel: +32.2.732.59.54.Fax: +32.2.732.60.03 START OF CE-MARKING: (Date or Lot or serial number) QINNGDAO. PLACE, DATE OF DECLARATION: Alex lin SIGNATURE: President

Appendix: list of (harmonised - EN) standards

| NO. | Reference | Title |
|-----|---|---|
| 1 | EN 60601-1:1990+ A1:1993+ A2:1995 | Medical electrical equipment - Part 1: General requirements for safety |
| 2 | EN 60601-1-2: 2007 (IEC60601-1-2:2007) | Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests |
| 3 | EN 60601-1-4:1996/A1: 1999 (IEC 60601-1-4:1996/A1:1999) | Medical electrical equipment - Part 1-4: General requirements for safety - Collateral standard: Programmable electrical medical systems |
| 4 | EN 60601-1-6:2007 (IEC 60601-1-6:2006) | Medical electrical equipment – Part 1-6: General requirements for safety- Collateral standard: usability |
| 5 | EN 60601-1-8: 2007 (IEC 60601-1-8: 2006) | Medical electrical equipment - Part 1-8: General requirements for safety - Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems |
| 6 | EN 60601-2-27:2006 (IEC 60601-2:27: 2005) | Medical electrical equipment - Part 2-27: Particular requirements for the safety, including essential performance, of electrocardiographic monitoring equipment |
| 7 | EN 60601-2-30:2000 (IEC 60601-2-30: 1999) | Medical electrical equipment - Part 2-30: Particular requirements for the safety, including essential performance, of automatic cycling non-invasive blood pressure monitoring equipment |
| 8 | EN 60601-2-34:2000 (IEC 60601-2-34: 2000) | Medical electrical equipment - Part 2-34: Particular requirements for the safety, including essential performance, of invasive blood pressure monitoring equipment |
| 9 | EN 60601-2-49: 2001 (IEC 60601-2-49: 2001) | Medical electrical equipment - Part 2-49: Particular requirements for the safety of multifunction patient monitoring equipment |
| 10 | EN 62304: 2006 (IEC 62304: 2006) | Medical device software - Software life-cycle processes |
| 11 | EN 1060-1:1995+A2:2009 | Non-invasive sphygmomanometers - Part 1: General Requirements |

| TF-CE081117.1-09 | Ver: E |
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| Page 2 of 3 | |

| 12 | EN 1060-3:1997+A2:2009 | Non-invasive sphygmomanometers - Part 3: Supplementary requirements for electro-mechanical blood pressure measuring systems |
|----|---|---|
| 13 | EN 12470-4:2000+A1:2009 | Clinical thermometers - Part 4: Performance of electrical thermometers for continuous measurement |
| 14 | EN ISO 9919: 2009 (ISO 9919: 2005) | Medical electrical equipment – Particular requirements for the basic safety and essential performance of pulse oximeters equipmet for medical use |
| 15 | EN ISO 21647:2009 (ISO 21647: 2004/ AC:2006) | Medical electrical equipment - Particular requirements for the basic safety and essential performance of respiratory gas monitors |