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

MANUFACTURER:

CONTEC MEDICAL SYSTEMS CO., LTD

PEOPLE'S REPUBLIC OF CHINA

No.112 Qinhuang West Street, Economic & Technical

Development Zone, Qinhuangdao, Hebei Province,

| MEDICAL DEVICE: | Electrocardiograph ECG600G | | | |
|--|--|--|--|--|
| CLASSIFICATION - ANNEX IX: | NEX 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 MEDICAL DEVICES. MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE 93/42/EEC OF 14 JUNE 1993 CONCERNING MEDICAL DEVICES; NCLUDING, 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: | TÜV SÜD PRODUCT SERVICE GMBH RIDLERSTR 65, D-80339 M NCHEN, GERMANY | | | |
| IDENTIFICATION NUMBER: | C € ₀₁₂₃ | | | |
| (EC) CERTIFICATE(S): | G1 050972 0050 Rev.04 | | | |
| EUROPEAN REPRESENTATIVE: | Shanghai International Holding Corp. GmbH(Europe) Eiffestrasse 80, 20537 Hamburg Germany | | | |
| START OF CE-MARKING: 2012-04-20 (Date or Lot or serial number) | | | | |
| PLACE, DATE OF DECLARATION: | QINHUANGDAO, 2020-06-18 | | | |
| SIGNATURE | President | | | |
| SIGNATURE: | rresidefit | | | |
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Appendix: list of (harmonised - EN) standards

| No. | Reference | Title of Standard |
|-----|---------------------|---|
| 1 | 150 00004 4 0040 | Medical electrical equipment - Part 1: General requirements for |
| | IEC 60601-1:2012 | basic safety and essential performance |
| 2 | | Medical electrical equipment - Part 1-2: General requirements for |
| | | basic safety and essential performance - Collateral Standard: |
| | | Electromagnetic disturbances - Requirements and tests |
| 3 | | Medical electrical equipment-Part 1-6:General requirements |
| | IEC 60601-1-6:2013 | for basic safety and essential performance-Collateral |
| | | Standard: Usability |
| | | Medical electrical equipment –Part 2-25: Particular |
| 4 | 120 00001 2 20.2011 | requirements for the basic safety and essential performance of |
| | | electrocardiographs |
| | 5 IEC 62366-1:2015 | Medical devices - Application of usability engineering to |
| 5 | | medical devices |
| 6 | IEC 62304:2015 | Medical device software-Software life-cycle processes |

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