



EC DECLARATION OF CONFORMITY

TF – DOC0077948 (CE-M-098)

Following the provisions of the medical device directive 93/42/EEC, Annex II and of the directive 2011/65/EU.

EG-KONFORMITÄTSERKLÄRUNG

Gemäß den Vorschriften der Richtlinie 93/42/EWG über Medizinprodukte, Anhang II und der Richtlinie 2011/65/EU.

We/ Wir

Manufacturer

Hersteller

**GE Medical Systems
Information Technologies, Inc.
8200 West Tower Avenue
Milwaukee, WI 53223, USA**

EU Authorized Representative

Autorisierter EU-Vertreter

**GE Medical Systems
Information Technologies GmbH
Munzinger Straße 5
79111 Freiburg, Germany**

Manufacturing site (if different from manufacturer)

Fertigungsstätte (falls anders als Hersteller)

**Critikon de Mexico S. de R.L. de C.V.
Calle Valle de Cedro 1551
Juarez, Chihuahua, Mexico 32575**

**GE Healthcare Finland Oy
Kuortaneenkatu 2
00510 Helsinki, Finland**

**GE Medical Systems Information Technologies
456 Pan American Drive, Suite 11
El Paso, Texas 79907, USA**

Declare under our sole responsibility that the class IIa medical device:

Erklären unter unserer alleinigen Verantwortung, dass das Medizinprodukt der Klasse IIa:

CardioSoft Cardiac Testing System

Including system components and accessories/einschließlich Systemkomponenten und Zubehör)

Ref. : see addendum/ oder siehe Anhang

GMDN Code: 36145

UMDNS Code: 17-723

Classification rule (93/42/EC Annex IX) / Klassifizierungsregel (93/42/EWG Anhang IX) : **Rule 10**

To which this declaration relates, is in conformity with the requirements of the medical device directive 93/42/EEC which apply to it and with the requirements of the directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment.

Auf das sich diese Erklärung bezieht, den Anforderungen der Richtlinie 93/42/EWG über Medizinprodukte, die für das Produkt gelten, und den Anforderungen der Richtlinie 2011/65/EU zur Beschränkung der Verwendung bestimmter gefährlicher Stoffe in Elektro- und Elektronikgeräten entspricht.


Lee Bush

Director, Regulatory Affairs

Milwaukee, USA, 12.April.2019

***This medical device conformity is based on the following elements:***

Diese Medizinprodukte Konformität basiert auf den folgenden Elementen:

- Information included in the documents:
Technical Documentation/DHF Ref./ réf: **DOC0410162**, of the product to which this declaration relates.
Informationen, die in den Dokumenten enthalten sind:
Technische Dokumentation/DHF-Ref./réf: **DOC0410162** des Produkts, auf das sich diese Erklärung bezieht.
- EC certificate: approval of full quality assurance system (Annex II of the medical devices directive 93/42 EEC) delivered by LNE/G-MED France, (NB #0459) Certificate No. 7550 ([DOC0279260](#)).
EG-Zertifikat: Genehmigung des kompletten Qualitätssicherungssystems (Anhang II der Richtlinie 93/42/EWG über Medizinprodukte), ausgestellt von G-MED France, NB #0459 / Zertifikat Nr. 7550 ([DOC0279260](#)).

List of harmonized standards applied for CE marking

Liste der harmonisierten Normen, die für die CE-Kennzeichnung angewendet wurden

1. **EN ISO 13485:2016** Medical devices -- Quality management systems -- Requirements for regulatory purposes
2. **EN ISO 14971:2012** Medical Devices – Application of Risk Management to Medical Devices
3. **EN 60601-1:1990, A1:1993, A2:1995, A13:1996**, Medical Electrical Equipment Part 1: General Requirements for Safety
4. **EN 60601-1:2012** Medical electrical equipment – Part 1: General Requirements for Basic Safety and Essential Performance
5. **EN 60601-1-1:2001** Medical electrical equipment – Part 1-1: General requirements for safety Collateral standard: Safety requirements for medical electrical systems
6. **EN 60601-1-2:2007/AC:2010** Medical electrical equipment – Part 1-2: General requirements for safety Collateral standard: Electromagnetic compatibility Requirements and tests
7. **EN 60601-1-2:2015** Medical electrical equipment – Part 1-2: General requirements for safety and essential performance – Collateral Standard: Electromagnetic disturbances – Requirements and tests
8. **EN 60601-1-4:1996, A1:1999** Medical Electrical Equipment – Part 1-4: General requirements for safety Collateral Standard: Programmable electrical medical systems
9. **EN 62304:2006** Medical device software-Software life-cycle processes
10. **EN 62366:2008** Medical devices - Application of usability engineering to medical devices
11. **EN 60601-1-6:2007** Medical electrical equipment - Part 1-6: General requirements for safety - Collateral standard: Usability
12. **EN 60601-2-25:1995, A1:1999** Medical Electrical Equipment – Part 2-25: Particular Requirements for the Safety of Electrocardiographs
13. **EN 60601-2-25:2011** Medical electrical equipment – Part 2-25: Particular requirements for the basic safety and essential performance of Electrocardiographs
14. **EN 60601-2-51:2003** Medical electrical equipment -- Part 2-51: Particular requirements for safety, including essential performance, of recording and analyzing single channel and multichannel electrocardiographs
15. **EN 1041:2008+A1:2013** Information supplied by the manufacturer with medical devices
16. **EN ISO 15223-1:2016** Symbols for use in the labeling of medical devices
17. **EN ISO 10993-1:2009/AC:2010** Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process
18. **EN ISO 10993-5:2009** Biological Evaluation of Medical Devices - Part 5: Test For In Vitro Cytotoxicity

This EC declaration of conformity supersedes the previous declaration dated 14-March-2019

Lee Bush

Director, Regulatory Affairs

Milwaukee, USA, 12.April.2019



Diese EG-Konformitätserklärung ersetzt die vorherige Erklärung mit Datum vom 14.März.2019

Milwaukee, USA, 12.April.2019

A handwritten signature in blue ink, appearing to read 'Lee Bush', written over a horizontal line.

Lee Bush
Director, Regulatory Affairs



ADDENDUM TO THE EC DECLARATION OF CONFORMITY
ERGÄNZUNG ZUR KONFORMITÄTSERKLÄRUNG

Product Description Produktbezeichnung	Catalog Designation Katalogbezeichnung
CardioSoft v6.7 CD	2006301-044
CardioSoftv6.7 Software	2006300-101
Options:	Part Number:
CardioSoft ECG Analysis System V6.7 Generic ATO Model	2060450-001
CSOFT-CS V6.7 CLIENT GENERIC ATO MODEL	2060452-001
CSOFT ECG VIEWER V6.7 GENERIC ATO MODEL	2060451-001
CAM-USB Interface Box V2 , required for the CAM-14 acquisition module	2040437-001
CAM-USB A/T Interface Box V2	2040438-001
CAM-USB A/T KISS Interface Box V2	2040438-009
Software Option:	
Resting ECG Interpretation (RESI)	45502401
Remote View (ERGM)	45502901
Storage of the Full-Disclosure ECG (EGMO)	45502701
Data Storage on Network Server, < 3000 tests (NETS)	45503001
Data Storage on Network Server, < 15000 tests (NET2)	2014659-001
Data Storage on Network Server, unlimited numbers of tests (NET3)	2014659-002
Arrhythmia Detection / Documentation (ARRY)	2014659-014
2D Waterfall Display (2DWF)	2014659-003
MUSE Browser (BRWS)	2014659-004
Risk Factors (RISK)	2014659-005
Data Export (EXPD)	2014659-006
Report Export as PDF File (EPDF)	2014659-007
Report Export as Word File (EWRD)	2014659-008
Display Configuration (DSPC)	2014659-009
In-Test Tabular Summary (ITBL)	2014659-010
In-Test Trend (ITRD)	2014659-011
Previous Test Retrieval (PRVT)	2014659-012
T-Wave Alternans (TWAA)	2014659-013
EMR Interface (XEMR)	2014659-020
Floating License (FLLX)	2014659-021
Resting ECG Measurement (RESM)	45502301
ST Measurement, Arrhythmia, 6/12-Lead Exercise Test (ERG2)	45502601
Exercise Test Expert Mode (ERG3)	45503201
CardioSoft Web (CWEB)	45505101
ECG History (ECGH)	45504001
DICOM interface (DICM)	2014659-026
ST/HR Hysteresis (STHY)	2014659-028
Exercise Test Interpretation (EXTI)	2014659-029

Lee Bush

Director, Regulatory Affairs

Milwaukee, USA, 12.April.2019