



Special 510(k) for
CG-6108 ACT-3L Continuous ECG monitor and Arrhythmia Detector
Section 7: 510(k) Summary

APR - 6 2011

510(k) Summary: Modified CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector

Introduction

This document contains the 510(k) summary for the modified CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector. The content of this summary is based on the requirements of 21 CFR Section 807.92(c).

Submitter	Card Guard Scientific Survival Ltd.,	
Establishment Registration Number	9681879	
Address	2 Pekeris St., P.O.B. 527, Rehovot, 76100, Israel	
Contact person:	Asher Kassel, Director of RA & QA, Card Guard Scientific Survival Ltd.	
Phone:	972-8-9484010 (direct)	Fax: 972-8-9484044
E-mail:	asher@cardguard.com	
Date Prepared:	February 18, 2011	
Predicate device	Unmodified version of CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector, cleared in K101703 on July 13, 2010.	
Trade Name:	CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector	
Classification:	Detector and alarm, arrhythmia /Transmitters and receivers, electrocardiograph, telephone	
Product Code:	DSI, DXH	
Regulation No:	870.1025, 870.2920	
Class:	II	

Device Description

The CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector is designed for self-testing by patients at home and for analysis by medical professionals at a remote Monitoring Center.

The chest-worn sensor is used for the acquisition, recording, and transmission of the ECG signal. The device is equipped with 4 electrodes on a harness and it houses a 3.6V AA battery, a Bluetooth transceiver and a buzzer.

The ECG signals are transmitted via Bluetooth to a handheld device with a proprietary interactive application, configured to process and transmit the ECG recordings. The handheld device is a mobile computing device with a display and a touch input such as a cell phone. It has sufficient memory and processing capability to run the proprietary application.

When an arrhythmia event is detected, the handheld device transmits the recorded ECG information automatically via cellular link, to the Monitoring Center for professional analysis. When cellular service is unavailable the patient has an option to transmit via a landline telephone.

The Patient and Physician manuals are being modified to change a warning to allow the use of the ACT-3L on patients with an Implanted Cardioverter Defibrillator (ICD) if specific precautions are observed.

Indications for Use

The CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector is intended for use by patients who experience transient symptoms that may suggest cardiac arrhythmia. The device continuously monitors patient ECG, automatically generates an alarm triggered by an arrhythmia detection algorithm, or generates an alarm manually triggered by the patient, and transmits the



recorded data transtelephonically to a monitoring center. The monitoring center provides the ECG data to the medical practitioner for evaluation.

Summary of the Technological Characteristics / Principles of Operation

The technological characteristics and principles of operation of the modified device are the same as the predicate device. The chest-worn ECG sensor transmits signals via Bluetooth to the handheld device equipped with the Medical Application, which incorporates an algorithm for detection of cardiac events: Atrial Fibrillation, Tachycardia, Bradycardia and Pause. A detected artifact triggers transmission of the signal to the Monitoring Center for analysis.

Non-clinical performance data for the CG-6108 ACT-3L

In order to support the labeling change the following testing has been performed:

- ACT-3L High Voltage Pulse Test, Card Guard document # ENTR-0112
- ACT-3L EMC Dipole Antenna Test, Card Guard document # ENTR-0113

Performance Standards:

This 510(k) submission was written in accordance with the FDA Guidance document "Class II Special Controls Guidance Document: Arrhythmia Detector and Alarm, October 28, 2003" and the device conforms to the applicable performance requirements contained in and referenced in this document. In addition, this submission was prepared in accordance with "Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices, May 11 2005". The design of the CG-6108 ACT-3L conforms to the following voluntary standards:

- ANSI/AAMI EC57:1998 (R) 2008: Testing and reporting performance results of cardiac rhythm and ST segment measurement algorithms (Cardiovascular)
- ANSI/AAMI EC38:1998 Medical electrical equipment – Part 2-47: Particular requirements for the safety, including essential performance, of ambulatory electrocardiographic systems (Cardiovascular)
- ISO 14971:2007: Medical devices – application of risk management to medical devices (General)
- IEC 60601-1:1988, 2nd edition, Part 1, plus A1:1991 and A2:1995: Medical electrical equipment; Part 1: General requirements for safety
- IEC 60601-1-2: 2001, plus A1:2004, Part 1: Medical electrical equipment, Part 1-2: General requirements for safety – Collateral standard: Electromagnetic compatibility - Requirements and tests
- IEC 60601-1-4:2000, Medical electrical equipment - Part 1:General requirements for safety; Part 1- 4: Collateral standard: Programmable electrical medical systems
- IEC 62304:2006: Medical device software – Software life cycle processes
- ISO 15223:2000: Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied – Part 1: General requirements

Substantial Equivalence:

The modified CG-6108 ACT-3L device is substantially equivalent with respect to the indications for use, technological characteristics and performance characteristics to the identified legally marketed predicate device.



Food and Drug Administration
10903 New Hampshire Avenue
Document Control Room - WO66-G609
Silver Spring, MD 20993-0002

Card Guard Scientific Survival Ltd
c/o Mr. Asher Kassel
Vice President of Operations
2 Pekeris St.
Rehovot, 76100 Israel

APR - 6 2011

Re: K110499

Trade/Device Name: CG-6108-3L Continuous ECG Monitor and Arrhythmia Detector
Regulation Number: 21 CFR 870.1025
Regulation Name: Arrhythmia detection or alarms (including ST-segment measurement and alarm)
Regulatory Class: Class II
Product Code: DSI and DXH
Dated: February 18, 2011
Received: February 22, 2011

Dear Mr. Kassel:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

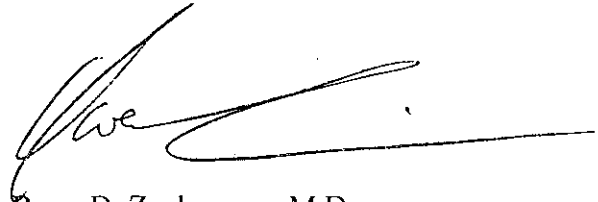
If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at 240-276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (240) 276-3150 or at its Internet address <http://www.fda.gov/cdrh/industry/support/index.html>.

Sincerely yours,



for

Bram D. Zuckerman, M.D.
Director
Division of Cardiovascular Devices
Office of Device Evaluation
Center for Devices and
Radiological Health

Enclosure

Indications for Use

510(k) Number (if known): K110499

Device Name: CG-6108 ACT-3L Continuous ECG Monitor and Arrhythmia Detector

Indications for Use:

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
Prescription Use X
(Part 21 CFR 801 Subpart D)

AND/OR

Over-The-Counter Use _____
(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of Device Evaluation (ODE)

for

(Division Sign-Off)
Division of Cardiovascular Devices

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