K063503

APR 27 2007

Special 510(k) Summary

1. Company Identification

Mallinckrodt Inc., Liebel-Flarsheim Business 2111 East Galbraith Road Cincinnati, OH 45237

Establishment Registration: 1518293

2. Contact Person

Dale Moore Quality Manager Phone: (513) 948-5771 Fax: (513) 948-5708 Email: dale.moore@tycohealthcare.com

3. 510(k) Preparation Date

11/15/2006

4. Device Name

Trade Name: OptiVantage DH Injector System with Enhanced Communication Common Name: Power Injector

5. Device Classification

Class II

6. Indications for Use

The OptiVantage DH Injector System with Enhanced Communication is a contrast delivery system and is designed to inject radiopaque contrast media into a patient's vascular system to obtain diagnostic images when used with computed tomography (i.e. "CT") equipment.

7. Description of Device

The OptiVantage DH Injector System with Enhanced Communication delivers radiographic contrast media and/or saline at a controlled flow rate and volume into a patient's vascular system for the purpose of obtaining enhanced diagnostic images. The OptiVantage DH Injector System with Enhanced Communication consists of 4 main components, just like the predicate device:

- 1. Power Head- Contains two electromechanical syringe drive systems, the syringe holding mechanisms, the main microprocessor, control electronics, control keypad for programming and initiating injection protocols, a status display, and two purge/retract manual knobs. The OptiVantage uses a drive system that is similar to existing Mallinckrodt injectors in the marketplace. It uses a motor and gearbox coupled to a ball screw. The ball screw drives a ram that attaches to the syringe plunger in order to fill or expel the contents of the syringe.
- 2. **Power Supply-** The power supply converts the line voltage to the working voltage for the power head and console (approximately 24-vdc).

- 3. Console Communicates with the Power Head to program and initiate injection protocols, displays the injection status, and displays a timer.
- 4. Syringes The OptiVantage DH Injector System with Enhanced Communication accommodates the Mallinckrodt 125-ml pre-filled syringe styles as well as a 200 ml front loading empty syringe. These syringes are commonplace on the market and are currently used with the predicate device.

8. Substantial Equivalence

The predicate injector to the OptiVantage DH Injector System with Enhanced Communication is the OptiVantage DH Power Injection System, 510(k) number K042744.

The OptiVantage DH Injector System with Enhanced Communication maintains the same intended use as the predicate device. It is intended to be used for the specific purpose of injecting radiopaque contrast media into a patient's vascular system to obtain diagnostic images when used with computed tomography (i.e. "CT") equipment.

The OptiVantage DH Injector System with Enhanced Communication consists of four main components in the same manner as the predicate device: a Power Head, a Console, a Power Supply, and Syringes. Both the OptiVantage DH Injector System with Enhanced Communication and the predicate device consist of the same fundamental technology. They are motor driven, electromechanical devices which are software controlled. The OptiBolus function on the OptiVantage DH Injector System with Enhanced Communication and the predicate device mplements a multiphasic injection method for injecting contrast media and/or saline.

The OptiVantage DH Injector System with Enhanced Communication differs from the predicate device in that it contains a product enhancement that adds a layer of enforcement for safety features found on the currently marketed predicate device.

Below is a table that compares the predicate device to the proposed OptiVantage DH Injector System with Enhanced Communication.

Feature	OptiVantage DH Injector System with	OptiVantage DH Power Injection
	Enhanced Communication (New Device)	System Predicate Device (K042744)
Intended Use	Intended to be used for the specific	Intended to be used for the specific
	purpose of injecting radiopaque contrast	purpose of injecting radiopaque contrast
	media into a patient's vascular system to	media into a patient's vascular system to
	obtain diagnostic images when used with	obtain diagnostic images when used with
	computed tomography (i.e. "CT")	computed tomography (i.e. "CT")
	equipment	equipment
Multi-phasic	6 phases per protocol	6 phases per protocol
Injections		
OptiBolus	Yes	Yes
Protocol Storage	40 protocols	40 protocols
Syringe Sizes	All pre-filled volumes of Mallinckrodt	All pre-filled volumes of Mallinckrodt
	125-ml syringes & Liebel-Flarsheim 200	125-ml syringes & Liebel-Flarsheim 200
	ml empty syringe	ml empty syringe
Single or Dual Syringe	Single syringe and Dual syringe capability	Single syringe and Dual syringe capability
Modality		
Syringe Drive System	Electromechanical	Electromechanical
Syringe Heater	Yes (Built-in, Powerhead controlled)	Yes (Built-in, Powerhead controlled)
Syringe Fill Rate	2 to 15 ml/sec	2 to 15 ml/sec
Flow Rate	0.1 to 10 ml/sec	0.1 to 10 ml/sec
Max Pressure Limit	325 psi	325 psi
Pressure Limit	User-settable or automatic	User-settable or automatic
Control	25 psi increments	25 psi increments
Flushing System	Manual or via injector	Manual or via injector

Feature	OptiVantage DH Injector System with Enhanced Communication (New Device)	OptiVantage DH Power Injection System Predicate Device (K042744)
Remote Start	Yes	Yes
Scan Delay	0-600 seconds	0-600 seconds
Phase Delays	0 – 600 seconds plus Pause	0 – 600 seconds plus Pause
Console Controls	Touch screen Color LCD	Touch screen Color LCD
Power head Controls	Touch screen Color LCD	Touch screen Color LCD
Post Injection Readout	Yes	Yes
OEM Interface	Relays & Optical Couplings/Serial (CAN)	Relays & Optical Couplings/Serial (CAN)
Safety Stop	Electrical Stop when injection parameters	Electrical Stop when injection parameters
Mechanism	are out of specifications	are out of specifications
Remote Check for Air	Yes	Yes
Volume Remaining	Displayed on Power Head and Console	Displayed on Power Head and Console
Display		
Used Syringe	Yes (with Enhanced Communication)	Yes
Detection		
Materials	Plastic and metal	Plastic and metal
Target Population	Humans	Humans
Sterility (Syringe)	Injectors are not sterile products. Syringes and Disposables are provided sterile.	Injectors are not sterile products. Syringes and Disposables are provided sterile.

Public Health Service



Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

APR 27 2007

Mallinckrodt, Inc., Liebel-Flarsheim Business c/o Mr. Dale Moore Quality Manager 2111 East Galbraith Road Cincinnati, OH 45237

Re: K063503

OptiVantage DH Injector System with Enhanced Communication Regulation Number: 21 CFR 870.1650 Regulation Name: Angiographic injector and syringe Regulatory Class: II (two) Product Code: IZQ Dated: March 30, 2007 Received: April 2, 2007

Dear Mr. Moore:

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.

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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" (21CFR 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 <u>http://www.fda.gov/cdrh/industry/support/index.html</u>.

Sincerely yours,

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

Enclosure

Healthcare Division

Attachment #14 Indications for Use

510(k) Number (if known): K063503

Device Name: OptiVantage DH Injector System with Enhanced Communication

Indications For Use:

The OptiVantage DH Injector System with Enhanced Communication is a contrast delivery system and is designed to inject radiopaque contrast media into a patient's vascular system to obtain diagnostic images when used with computed tomography (i.e. "CT") equipment.

Prescription Use X (Part 21 CFR 801 Subpart D) AND/OR

Over-The-Counter Use _____ (21 CFR 807 Subpart C)

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

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)

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(Division Sign-Off) Division of Cardiovascular Devices

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