

FEB - 3 2012

9. 510(K) SUMMARY

This summary of 510(k) safety and effectiveness information is submitted in accordance with the requirements of SMDA 1990 and 21 CFR 807.92.

510(k) Number: K120129

Applicant Information:

Date Prepared: January 13, 2012

Name: BridgePoint Medical
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Plymouth, MN 55441
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Contact Person: Jill Munsinger
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E-mail: jmunsinger@bridgepointmedical.com

Device Information:

Classification: Class II Percutaneous Catheter
Trade Name: Mantaray™ Catheter
Common Name: Percutaneous Catheter
Classification Name: Percutaneous Catheter

Predicate Devices:

The modified BridgePoint Medical Mantaray™ Catheter is substantially equivalent in intended use, method of operation and technical aspects to the following predicate device:

K111963 - Mantaray™ Catheter

Device Description:

The modified Mantaray™ Catheter is a single use, over-the-wire, disposable, dual lumen percutaneous catheter that facilitates the placement, support and steering of a guidewire into discrete regions of the peripheral vasculature through the central guidewire lumen or through one of two sideports (identified by radiopaque markers). The sideports connect with the central guidewire lumen and facilitate guidewire steering (at an angle to the central lumen) by allowing the guidewire to exit the catheter. The catheter contains a small non-compliant balloon segment used for fluoroscopic orientation on the distal tip of the flexible shaft.

Intended Use:

The BridgePoint Medical Mantaray™ Catheter is indicated for directing, steering, controlling, and supporting a guidewire in order to access discrete regions of the peripheral vasculature.

Comparison to Predicate Device(s):

The modified Mantaray™ Catheters are substantially equivalent to the current Mantaray™ Catheters, K111963 in that they are both designed to direct, steer, control and support a guidewire in accessing discrete regions of the peripheral vasculature.

The modified Mantaray™ Catheter is constructed of the same materials as the current Mantaray™ Catheter. Modifications were made to remove the coating on the balloon segment, which alleviates the need to soak the device for 10 minutes prior to use. The modified and current Mantaray™ Catheters are manufactured using similar processes and components and have similar physical attributes (balloon performance, trackability, tensile, radiopacity, and torque, etc.).

Both devices include radiopaque markers located within the balloon segment to indicate the location of the guidewire lumen ports.

Performance Data:

The modified Mantaray™ Catheters have been evaluated using the following *in vitro* bench testing to confirm the performance characteristics as compared to the predicate device:

- Tensile
- Burst
- Fatigue
- Inflation & Deflation Time
- Dimensional
- Hydration
- Guidewire Insert & Withdrawal
- Flexibility
- Trackability
- Guidewire Re-Direction
- Markerband Movement & Removal
- Markerband & Guidewire Interaction
- Kink Resistance
- Coating
- Torque
- Surface Defects
- Balloon Protector Removal, and

In vivo testing was not deemed necessary based on the significance of the proposed modification to the baseline device. Removal of the coating from the distal 6 cm of the catheter results in a device that meets the original design requirements of the currently

marketed Mantaray™ Catheter as demonstrated in the bench tests above. Animal studies were successfully completed with the currently marketed Mantaray™.

Likewise, the removal of coating from the distal 6 cm of the device did not warrant additional biocompatibility testing as all materials are included in the currently marketed Mantaray™ Catheter.

Summary:

Based upon the intended use and descriptive information provided in this pre-market notification, the modified BridgePoint Mantaray™ Catheters have been shown to be substantially equivalent to the currently marketed predicate device.



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

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BridgePoint Medical, Inc.
c/o Jill Munsinger
13355 10th Ave N
Plymouth, MN 55441

Re: K120129
Trade/Device Name: Mantaray™ Catheter
Regulation Number: 21 CFR 870.1250
Regulation Name: Percutaneous Catheter
Regulatory Class: Class II
Product Code: DQY
Dated: January 13, 2012
Received: January 17, 2012

Dear Ms. Munsinger:

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. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you; however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to 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); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); 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.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please go to <http://www.fda.gov/AboutFDA/CentersOffices/CDRH/CDRHOffices/ucm115809.htm> for the Center for Devices and Radiological Health's (CDRH's) Office of Compliance. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

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 (301) 796-7100 or at its Internet address <http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,



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

Enclosure

8. INDICATIONS FOR USE STATEMENT

510(k) Number: (TBA)

Device Name: BridgePoint Medical Mantaray™ Catheter

Indications For Use:

The BridgePoint Medical Mantaray™ Catheter is indicated for directing, steering, controlling, and supporting a guidewire in order to access discrete regions of the peripheral vasculature.

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 Device Evaluation (ODE)

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(Division Sign-Off)
Division of Cardiovascular Devices

510(k) Number K120129