5. 510(k) Summary

Date Prepared:
February 28, 2012

Submitter's Information:
FUJIFILM Medical Systems U.S.A., Inc.
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Stamford, Connecticut 06902

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Contact: Jyh-Shyan Lin

Device Trade Name:
Synapse 3D Cardiac Tools

Device Common Name:
Medical Image Processing and Analysis Software

Regulation Number:
21 CFR 892.2050

Device Classification:
Class II

Device Classification Name:
Picture Archiving and Communications System (PACS)

Panel:
Radiology

Product Code:
LLZ

Date Received:
TBD

Decision Date:
TBD

Decision:
TBD
Predicate Device:

- Synapse 3D Cardiac Tools, FUJIFILM Medical Systems U.S.A., Inc. (K103465)
- QPlaque MR, Medis Medical Imaging Systems, B.V. (K073156)

Description of the Device

Synapse 3D Cardiac Tools (V3.0) is the updated version of previously-cleared Synapse 3D Cardiac Tools software (cleared by CDRH via K103465 on 01/31/2011).

Synapse 3D Cardiac Tools is used in addition to the Synapse 3D Base Tools (K120361, cleared on April 6, 2012) to analyze the images acquired from CT and MR. Synapse 3D Cardiac Tools is intended to provide trained medical professionals with tools to aid them in reading, interpreting, reporting, and treatment planning of DICOM compliant medical images. This product is not intended for use with or for the primary diagnostic interpretation of mammography images.

Synapse 3D Cardiac Tools (V3.0) is an application that supports the cardiac function, cardiac fusion, and coronary artery analysis of both the computed tomography (CT) and magnetic resonance (MR) images. Synapse 3D Cardiac Tools also supports the calcium scoring for non-contrast CT images.

Unchanged Applications from the cleared version K103465

(1) Cardiac Function (CT)
   Cardiac Function (CT) is an application for cardiac function evaluation which obtains the boundary between left ventricle and cardiac wall from CT left ventriculography images retrieved from multiple time phases and calculates ejection fraction, end-diastolic volume, end-systolic volume, stroke volume, etc.

(2) Cardiac Function (MR)
   Cardiac Function (MR) is an application for cardiac function evaluation which obtains the boundary between left ventricle and cardiac wall from non-contrast MR images retrieved from multiple time phases and calculates ejection fraction, end-diastolic volume, end-systolic volume, output volume per beat, etc.

(3) Coronary Artery Analysis (CT)
   Coronary Artery Analysis (CT) is an application using CT coronary arteriography images to extract the path of the target blood vessels and to perform coronary artery evaluation.

(4) Calcium Scoring
   The calcium scoring is an application which uses non-contrast CT images to display the calcification area in the coronary artery with color separation and calculates the calcification quantitative values using the Agatston score method.

(5) Cardiac fusion
   Cardiac fusion is an application to create an image having the mutual characteristics of source images of heart. Source images could be original image of CT, MR or NM and the functional image derived from the original image.
New Application

(6) Coronary Artery Analysis (MR)

Coronary Artery Analysis (MR) is an application using contrast or non-contrast MR heart images to extract the path of the target blood vessels and to perform coronary artery evaluation.

Note: The detail features available in this application are very similar to those in the Coronary Artery Analysis (CT).

In addition to the common image processing functions (such as window width and window level, zooming, panning, flip, rotation, adding annotations on an image, measurement of lengths, areas, etc.), the following image processing tools are available to support the cardiac analysis of the CT and MR images. These tools belong to and are provided by Synapse 3D Base Tools (K120361) that is used with Synapse 3D Cardiac Tools (V3.0).

- SUV evaluation: SUV average, standard deviation, etc. can be measured.
- Extraction and Deletion of 3D objects: Editing of mask areas using the smart cut feature.
- 3D clipping: The display area can be specified for 3D display.
- Organ segmentation and removal: Organs and other areas of interest in the image data can be segmented or removed.
- Mask editing: The mask area can be edited by lines drawn in freehand.
- CPR: CPR images can be created along a specified center line.
- Reformat: Plane images in any direction can be created.
- Creation of video files: Video files with 2D or 3D display can be created.
- Surface display: A polygon model of an image can be created.

Indication for Use

Synapse 3D Cardiac Tools is medical imaging software used with Synapse 3D Base Tools that is intended to provide trained medical professionals with tools to aid them in reading, interpreting, reporting, and treatment planning. Synapse 3D Cardiac Tools accepts DICOM compliant medical images acquired from a variety of imaging devices including, CT, MR, NM, and XA.

This product is not intended for use with or for the primary diagnostic interpretation of Mammography images.

Addition to the tools in Synapse 3D Base Tools, Synapse 3D Cardiac Tools provides the tools for specific clinical applications which provide targeted workflows, custom UI, targeted measurements and reporting functions including:

- Functional cardiac analysis for CT left ventriculography images: which is intended to evaluate the functional characteristics of heart
- Functional cardiac analysis for non-contrast MR heart images: which is intended to evaluate the functional characteristics of heart
Coronary artery analysis for CT coronary arteriography images: which is intended for the qualitative and quantitative analysis of coronary arteries

Coronary artery analysis for MR heart images: which is intended for the qualitative and quantitative analysis of coronary arteries

Calcium scoring for non-contrast CT heart images: which is intended for non-invasive identification and quantification of calcified atherosclerotic plaques in the coronary arteries using tomographic medical image data and clinically accepted calcium scoring algorithms

Cardiac Fusion: which is intended to analyze cardiac anatomy and pathology with a fused image of functional data (e.g. NM image, Bulls eye) and anatomical data.

Technological Characteristics

Synapse 3D Cardiac Tools introduces no new safety or efficacy issues other than those already identified with the predicate devices. The results of the Hazard Analysis combined with the appropriate preventive measures taken indicate that the device is of moderate concern as per the May 11, 2005 issue of the "Guidance for the Content of Premarket Submission for Software Contained in Medical Devices."

Testing

Synapse 3D Cardiac Tools is tested successfully with reference to its Software Requirements Specification, as well as design verification and validation documents and Traceability Matrix document. Verification, validation, and testing activities establish the performance, functionality, and reliability characteristics of the Synapse 3D Cardiac Tools software, which is found to be safe and effective and substantially equivalent to the currently-cleared predicate device.

Conclusion

This 510(k) premarket notification submission has demonstrated Substantial Equivalence as defined and understood in the Federal Food Drug and Cosmetic Act and various guidance documents issued by the Center for Devices and Radiological Health. We conclude the subject device to be as safe and effective as the predicate devices.
FUJIFILM Medical Systems U.S.A., Inc.
C/O JY-SHYAN LIN, Ph.D.
Senior Manager, Regulatory, Quality and Clinical Affairs
419 West Avenue
Stamford, Connecticut 06902

Re: K120636
Trade/Device Name: Synapse 3D Cardiac Tools
Regulation Number: 21 CFR 892.2050
Regulation Name: Picture Archiving and Communications System (PACS)
Regulatory Class: Class II
Product Code: LLZ
Dated: June 21, 2012
Received: June 22, 2012

Dear Mr. Lin:

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 class II (Special Controls), it may be subject to such additional controls. Existing major regulations affecting your device can be found in Title 21, Code of Federal Regulations (CFR), Parts 800 to 895. 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 Parts 801 and 809); medical device reporting (reporting of
medical device-related adverse events) (21 CFR 803); and good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820). 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 Parts 801 and 809), please contact the Office of In Vitro Diagnostic Device Evaluation and Safety at (301) 796-5450. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR 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/cdrh/industry/support/index.html.

Sincerely Yours,

Janine M. Morris
Aging Director
Division of Radiological Devices
Office of In Vitro Diagnostic Device Evaluation and Safety
Center for Devices and Radiological Health

Enclosure
Indications for Use

510(k) Number (if known):

Device Name: Synapse 3D Cardiac Tools

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- Cardiac Fusion: which is intended to analyze cardiac anatomy and pathology with a fused image of functional data (e.g. NM image, Bulls eye) and anatomical data.

Prescription Use X AND/OR Over-The-Counter Use

(Part 21 CFR 801 Subpart D) (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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