

510(k) Summary: syngo DynaPBV Body Software

JUL 30 2012

Company: Siemens Medical Systems, Inc.
51 Valley Stream Parkway
Malvern, PA 19355

Date Prepared: July 25, 2012

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

1. General Information:**Importer / Distributor:**

Siemens Medical Systems, Inc.
51 Valley Stream Parkway
Malvern, PA 19355

Establishment Registration Number:

2240869

Manufacturing Site:

SIEMENS AG Sector Healthcare
Siemensstraße 1
D-91301 Forchheim, Germany

Establishment Registration Number:

3004977335

2. Contact Person:

Ms. Patricia D Jones
Technical Specialist, Regulatory Submissions
Siemens Medical Solutions USA, Inc.
51 Valley Stream Parkway D-02
Malvern, PA 19355
Phone: (610) 448 -3536 Fax: (610) 448-1787
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3. Device Name and Classification:

Trade Name: syngo DynaPBV Body
Classification Name: System, Image Processing Radiological
Classification Panel: Radiology
Classification Regulation: 21 CFR §892.2050
Device Class: Class II
Product Code: LLZ

4. Legally Marketed Predicate Device

Trade Name: InSpace 3D Software Option
510(k) #: K011447
Clearance Date: August 3, 2001

Classification Name: Accessory to Angiographic X-Ray System
Classification Panel: Radiology
CFR Section: 21 CFR §892.1600
Device Class: Class II
Product Code: JAA

Legally Marketed Predicate Device

Trade Name: syngo Neuro PBV
510(k) #: K111052
Clearance Date: May 20, 2011
Classification Name: System, Image Processing Radiological
Classification Panel: Radiology
CFR Section: 21 CFR §892.2050
Device Class: Class II
Product Code: LLZ

5. Device Description:

The *syngo* DynaPBV Body software is an optional extension to the InSpace 3D application originally cleared under Premarket Notification K011447 on 08/03/2001. It is also similar to the cleared *syngo* Neuro PBV IR (K111052, May 20, 2011) which was designed for the visualization of contrast enhanced blood distribution in the arterial and venous vessels in the head.

Similar to *syngo* Neuro PBV IR the *syngo* DynaPBV Body is an add-on software option used for the visualization of contrast enhanced blood distribution in the body (e.g. thorax and abdomen) using color coded relative values for diagnosis.

This software modification does not affect the intended use of the device nor does it alter its fundamental scientific technology.

6. Indication for Use:

The *syngo* DynaPBV Body is an extended software application to the InSpace 3D software option which allows the reconstruction of two-dimensional images acquired with a standard angiographic C-arm device into a three-dimensional image format.

The *syngo* DynaPBV Body is intended for imaging primarily soft tissue for diagnosis, surgical planning, interventional procedures and treatment follow-up. It is design for the visualization of contrast enhanced blood distribution in the body using color coded relative values for diagnosis.

This software is designed to visually assist physicians in the diagnosis and treatment of vessel malformations (i.e. Aneurysms, AVM's and Stenoses).

7. Substantial Equivalence:

The *syngo* DynaPBV Body software application is substantially equivalent to the commercially available Siemens software application, *syngo* Neuro PBV IR which is an option to the Siemens InSpace 3D software. The InSpace 3D software option was described in premarket notification K011447 which received 510(k) clearance

on August 03, 2001. The *syngo* Neuro PBV IR software option was described in premarket notification K111052 and received 510(k) clearance on May 20, 2011.

The *syngo* DynaPBV Body software is an optional extension to InSpace 3D and uses the same hardware and software components as the InSpace 3D software.

8. Summary of Technological Characteristics of the Subject Device as Compared with the Predicate Device:

The *syngo* DynaPBV Body is a software extension to InSpace 3D. The *syngo* DynaPBV Body features the same post processing software, user interface, archiving and communication as the predicate InSpace 3D. The *syngo* DynaPBV Body software is not a stand-alone software. It interfaces with InSpace 3D. The *syngo* DynaPBV Body user function keys are integrated into the InSpace 3D task card. The user function is similar to InSpace 3D task card except for an additional activation button for the *syngo* DynaPBV Body software features.

9. General Safety and Effectiveness Concerns:

Instructions for use are included within the device labeling, and the information provided will enable the user to operate the device in a safe and effective manner.

Risk management is ensured via a hazard analysis, which is used to identify potential hazards. These potential hazards are controlled via software development, verification and validation testing. To minimize electrical, mechanical and radiation hazards, Siemens adheres to recognized and established industry practice, and all equipment is subject to final performance testing. Furthermore, the operators are health care professionals familiar with and responsible for the evaluating and post processing of X-ray images.

10. Conclusion as to Substantial Equivalence:

syngo DynaPBV Body software is intended for similar indications as cleared in the predicate InSpace 3D. The *syngo* DynaPBV Body Software add-on application is designed for use with the InSpace 3D (K011447) for the visualization of contrast enhanced blood distribution in the body using color coded relative values for diagnosis.

The functionality of *syngo* DynaPBV Body software is similar to the predicate device. It is Siemens opinion, that the *syngo* DynaPBV Body add-on software is substantially equivalent to the InSpace 3D software (K011447) and the *syngo* Neuro PBV IR software (K111052).



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

JUL 30 2012

Ms. Patricia D. Jones
Technical Specialist, Regulatory Submissions
Siemens Medical Solutions USA, Inc.
51 Valley Stream Parkway D-02
MALVERN PA 19355

Re: K121292

Trade/Device Name: *Syngo DynaPBV Body*
Regulation Number: 21 CFR 892.2050
Regulation Name: Picture archiving and communications system
Regulatory Class: II
Product Code: LLZ
Dated: July 25, 2012
Received: July 26, 2012

Dear Ms. Jones:

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.

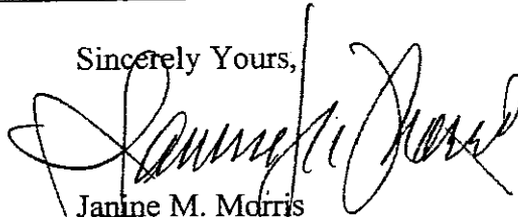
Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's 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
Acting Director
Division of Radiological Devices
Office of In Vitro Diagnostic Device
Evaluation and Safety
Center for Devices and Radiological Health

Enclosure

Indications for Use Statement

510(k) Number (if known): _____

Device Name: *syngo DynaPBV Body*

Indications for Use:

syngo DynaPBV Body is an extended software application to the InSpace 3D software option which allows the reconstruction of two-dimensional images acquired with a standard angiographic C-arm device into a three-dimensional image format.

syngo DynaPBV Body is intended for imaging primarily soft tissue for diagnosis, surgical planning, interventional procedures and treatment follow-up. It is design for the visualization of contrast enhanced blood distribution in the body using color coded relative values for diagnosis.

This software is designed to visually assist physicians in the diagnosis and treatment of vessel malformations (i.e. Aneurysms, AVM's and Stenoses)

Prescription Use X AND/OR Over-The-Counter Use _____
(Part 21 CFR 801 Subpart D) (21 CFR 801 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)



Division Sign-Off
Office of In Vitro Diagnostic Device Evaluation
and Safety

510(k) K121292

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