August 2, 2018

Philips Medical Systems Nederland BV
% Mr. Owen Callaghan
Regulatory Affairs Manager
Veenpluis 4-6
Best, 5684PC
NETHERLANDS

Re: K181830
  Trade/Device Name: Azurion R2.0
  Regulation Number: 21 CFR 892.1650
  Regulation Name: Image-intensified fluoroscopic x-ray system
  Regulatory Class: II
  Product Code: OWB, JAA
  Dated: July 6, 2018
  Received: July 9, 2018

Dear Mr. Callaghan:

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. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm identifies combination product submissions. 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.

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) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/CombinationProducts/GuidanceRegulatoryInformation/ucm597488.htm); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

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 comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/) and CDRH Learn (http://www.fda.gov/Training/CDRHLearn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (http://www.fda.gov/DICE) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

[Signature]

Robert Ochs, Ph.D.
Director
Division of Radiological Health
Office of In Vitro Diagnostics and Radiological Health
Center for Devices and Radiological Health

Enclosure
Indications for Use

510(k) Number (if known)
K181830

Device Name
Azurion R2.0

Indications for Use (Describe)
The Azurion series (within the limits of the used Operation Room table) are intended for use to perform:

- Image guidance in diagnostic, interventional and minimally invasive surgery procedures for the following clinical application areas: vascular, non-vascular, cardiovascular and neuro procedures.
- Cardiac imaging applications including diagnostics, interventional and minimally invasive surgery procedures.

Additionally:
- The Azurion series can be used in a hybrid Operation Room.
- The Azurion series contains a number of features to support a flexible and patient centric procedural workflow.

Patient Population
All human patients of all ages. Patient weight is limited to the specification of the patient table.

Type of Use (Select one or both, as applicable)
- [x] Prescription Use (Part 21 CFR 801 Subpart D)
- [ ] Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

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510(k) Summary

This 510(k) summary of safety and effectiveness information is prepared in accordance with 21 CFR §807.92.

Date Prepared: July 6th 2018

Manufacturer: Philips Medical Systems Nederland B.V.
Veenpluis 4-6
5684 PC Best
The Netherlands
Establishment Registration Number: 3003768277

Primary Contact Person: Owen Callaghan
Regulatory Affairs Manager
Phone: +31 621394159
E-mail: owen.callaghan@philips.com

Secondary Contact Person: Marta Walker
Head of Regulatory Affairs IGT Systems
Phone: +31 631978546
E-mail: marta.walker@philips.com

Device: Trade Name: Azurion R2.0
Classification Name: Image-intensified fluoroscopic x-ray system
Classification Regulation: 21 CFR, Part 892.1650
Classification Panel: Radiology
Device Class: Class II
Product Code: Primary Code: OWB
Secondary product code: JAA

Predicate Device: Trade Name: Azurion series R1.2
Manufacturer: Philips Medical Systems Nederland B.V.
510(k) Clearance: K172822 (November 22, 2017)
Classification Name: Image-intensified fluoroscopic x-ray system
Classification Regulation: 21 CFR, Part 892.1650
Classification Panel: Radiology
Device Class: Class II
Product Code: Primary Code: OWB
Secondary product code: JAA
Device description: The Azurion R2.0 is classified as an interventional fluoroscopic X-ray system. The primary performance characteristics of the Azurion R2.0 interventional fluoroscopic X-ray system include:

- Real-time image visualization of patient anatomy during procedures
- Imaging techniques and tools to assist interventional procedures
- Post processing functions after interventional procedures
- Storage of reference/control images for patient records
- Compatibility to images of other modalities via DICOM
- Built in radiation safety controls

This array of functions offers the physician the imaging information needed to perform minimally invasive interventional procedures.

The Azurion R2.0 is available as a single monoplane (single C-arm) configuration of the currently marketed and predicate devices Azurion series R1.2 consisting of a FD20 (frontal channel) detector with a ceiling (Clea) geometry having a standard or OR compatible table. This can optionally be configured with a horizontal L-arm which provides an extra rotation axis, with re-designed motion control software providing patient and X-ray beam positioning movement and an improved 3D roll scan time.

Additionally, identical to the predicate devices, Azurion R2.0 can be used in a hybrid operating room when supplied with a compatible operating room table, and can be optionally equipped with the ClarityIQ image processing algorithms.

Indications for Use: The Azurion series (within the limits of the used Operating Room table) are intended for use to perform:

- Image guidance in diagnostic, interventional and minimally invasive surgery procedures for the following clinical application areas: vascular, non-vascular, cardiovascular and neuro procedures.
- Cardiac imaging applications including diagnostics, interventional and minimally invasive surgery procedures.

Additionally:

- The Azurion series can be used in a hybrid Operating Room.
- The Azurion series contain a number of features to support a flexible and patient centric procedural workflow.

Patient Population:

All human patients of all ages. Patient weight is limited to the specification of the patient table.
Based on the information provided above, the **Azurion R2.0** is considered substantially equivalent to the currently marketed and predicate device **Azurion series R1.2** in terms of Indications for Use.

**Technological characteristics:**

The **Azurion R2.0** has similar technological characteristics compared to the predicate device. The same hardware and software is used in the predicate and subject device, with exception of the following modifications implemented in the **Azurion R2.0**:

- Introduction of:
  - Modified Horizontal L-arm with additional rotational axis
  - Rotatable Collimator
  - New Roll Drive
  - New motion control software for patient and X-ray beam positioning, giving an improved roll scan time
  - Automatic Image Beam Rotation
  - 2D image overlay on 3D model for additional C-arm positions (±45º and ±135º)
  - Additional monitor ceiling suspension boom

- Re-design of X-ray tube, Collimator
- Removal of X-ray tube side BodyGuard, replace with extended (3D) intelligent collision protection
- More consistent BodyGuard collision prevention (software update)
- Update of software to maintain Maquet table compatibility

The differences between the **Azurion R2.0** and the predicate device do not raise any new questions regarding safety or effectiveness. Based on the information provided in this 510(k) submission, **Azurion R2.0** is considered substantially equivalent to the currently marketed predicate **Azurion series R1.2** in terms of fundamental scientific technology.

**Summary of Non-Clinical Performance Data:**

Non-clinical performance testing has been performed on the **Azurion R2.0** and demonstrates compliance with the following International and FDA-recognized consensus standards and FDA guidance documents:


Software verification testing of the functional and non-functional requirements as well as performance, reliability and safety has been performed to verify that all
the requirements of System Requirements Specification as well as the safety risk control measures from the Detailed Risk Management Matrix and the Privacy and Security requirements have been implemented. Results demonstrated that all executed verification tests were passed.

Non-clinical validation testing has been performed to validate that Azurion R2.0 conforms to the intended use, claims, user and service needs, effectiveness of safety measures and instructions for use. The validation consisted of the following activities:

Usability validation was performed with both interventional radiologists/cardiologists (physicians) and nurse/technicians in a simulated use environment. Azurion R2.0 was found to be safe and effective for the intended use, users and use environment.

A simulated use design validation was performed with users that fulfill the intended user profile. The participants executed validation protocols in the form of a clinical workflow to validate user needs, intended use, claims and effectiveness of the safety and instructions for use. Results demonstrated that all executed validation protocols were passed.

Additionally, a simulated use study was conducted with interventional physicians to substantiate commercial product claims.

All these tests were used to support substantial equivalence of the subject device and demonstrate that Azurion R2.0:

- complies with the aforementioned international and FDA-recognized consensus standards and FDA guidance documents, and
- meets the acceptance criteria and is adequate for its intended use.

Therefore, Azurion R2.0 is substantially equivalent to the currently marketed Azurion series R1.2 in terms of safety and effectiveness.

Summary of Clinical Performance Data: The Azurion series R2.0 did not require clinical study data since substantial equivalence to the currently marketed predicate device Azurion series R1.2 was demonstrated with the following attributes:

- Indication for use;
- Technological characteristics;
- Non-clinical performance testing; and
- Safety and effectiveness.

These attributes demonstrated that the clinical performance of the modified device is substantially equivalent to the predicate device.
The **Azurion R2.0** is substantially equivalent to the currently marketed predicate device *Azurion series R1.2* in terms of indications for use, technological characteristics and safety and effectiveness.

The modification of the **Azurion R2.0** is within the controls and predetermined specifications. Additionally, substantial equivalence was demonstrated by non-clinical performance tests provided in this 510(k) premarket notification. These tests demonstrate that **Azurion R2.0** complies with the user need requirements as well as the requirements specified in the FDA-recognized consensus standards and guidance documents.

Therefore **Azurion R2.0** is as safe and effective as its predicate device and does not raise any new safety and/or effectiveness concerns.