



June 25, 2021

Boston Scientific Corporation  
Rochell Aranha  
Regulatory Affairs Specialist I  
Three Scimed Place  
Maple Grove Hennepin, Minnesota 55311

Re: K210889

Trade/Device Name: iLab Polaris Multi-Modality Guidance System  
Regulation Number: 21 CFR 870.1425  
Regulation Name: Programmable Diagnostic Computer  
Regulatory Class: Class II  
Product Code: DQK, DSK, ITX, IYO  
Dated: March 24, 2021  
Received: March 25, 2021

Dear Rochell Aranha:

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/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products>); 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 <https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance>) and CDRH Learn (<https://www.fda.gov/training-and-continuing-education/cdrh-learn>). 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 (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice>) for more information or contact DICE by email ([DICE@fda.hhs.gov](mailto:DICE@fda.hhs.gov)) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

LT Stephen Browning  
Assistant Director  
Division of Cardiac Electrophysiology, Diagnostics  
and Monitoring Devices  
Office of Cardiovascular Devices  
Office of Product Evaluation and Quality  
Center for Devices and Radiological Health

Enclosure

## Indications for Use

510(k) Number (if known)

K210889

Device Name

iLab™ Polaris Multi-Modality Guidance System

Indications for Use (Describe)

The IVUS modality of the iLab™ Polaris Multi-Modality Guidance System is intended for ultrasound examinations of intravascular pathology. Intravascular ultrasound is indicated in patients who are candidates for transluminal interventional procedures such as angioplasty and atherectomy.

FFR and DFR™ are intended for use in catheterization and related cardiovascular specialty laboratories to compute, and display various physiological parameters based on the output from one or more electrodes, transducers, or measuring devices. FFR and DFR are indicated to provide hemodynamic information for use in the diagnosis and treatment of patients that undergo measurement of physiological parameters.

The Imaging Catheters generate ultrasound images and are intended for ultrasound examination of vascular and cardiac pathology. Boston Scientific manufactures a wide variety of catheters for different applications. The recommended use of each of these catheters may vary depending on the size and type of the catheter. Please refer to the Imaging Catheter Directions for Use, packaged with each catheter.

Indications for Auto Pullback Use (IVUS Only)

Automatic Pullback is indicated when the following occurs:

- The physician/operator wants to standardize the method in which intravascular ultrasound images are obtained and documented: procedure-to-procedure, operator-to-operator.
- The physician/operator wants to make linear distance determinations post-procedurally, which requires the imaging core of a catheter to be pulled back at a known uniform speed.
- Two-dimensional, longitudinal reconstruction of the anatomy is desired.

Type of Use (Select one or both, as applicable)

Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

**CONTINUE ON A SEPARATE PAGE IF NEEDED.**

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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## 510(k) Summary per

### 21 CFR 807.92

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<b>Sponsor</b>	Boston Scientific Corporation 300 Boston Scientific Way Marlborough, Massachusetts 01752
<b>Contact Name and Information</b>	Rochell Aranha Specialist I, Regulatory Affairs Phone: 763-255-0017 Fax: 763-257-6482 e-mail: Rochell.Aranha@bsci.com
<b>Date Prepared</b>	March 24, 2021
<b>Proprietary Name</b>	iLab™ Polaris Multi-Modality Guidance System
<b>Common Name</b>	Computer Diagnostic Programmable
<b>Product Code</b>	DQK, DSK, IYO, ITX
<b>Classification</b>	Class II, 21 CFR 870.1425
<b>Predicate Name</b>	iLab™ Polaris Multi-Modality Guidance System K201178, May 29, 2020

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#### Device Description

The iLab™ Polaris Multi-Modality Guidance System is a non-patient contacting, diagnostic device designed to provide three (3) primary modalities: IVUS (Intravascular Ultrasound), FFR (Fractional Flow Reserve) and Diastolic hyperemia-Free Ratio (DFR). The iLab Polaris Multi-Modality Guidance System is supported by the Polaris 2.14 (iLab 3.14) software.

The iLab™ Polaris Multi-Modality Guidance System is offered in both mobile (cart) and integrated (installed) configurations consisting of hardware and software components supporting intravascular ultrasound (IVUS), fractional flow reserve (FFR) and Diastolic hyperemia-Free Ratio™ (DFR™) functionalities and includes additional Auto Pullback options.

Hardware components of the iLab™ Polaris Multi-Modality Guidance System include: Data Acquisition Processor, Imaging Processor, Control Panel Primary Image display (LCD flat panel), System Power supply/transformer, updated Motor Drive Unit (MDU5+), Motor Drive cable storage, Printer, removable Storage Media, CD/DVD Recorder, Mouse/Mouse Pad, and Bluetooth Communication Module (BCM).

The iLab Polaris Multi-Modality Guidance System includes automatic pullback functionality to aid the physician/operator during IVUS modality when the following occur:

- the physician/operator wants to standardize the method in which intravascular ultrasound images are obtained and documented: procedure-to-procedure, operator-to-operator,
  - the physician/operator wants to make linear distance determinations post-procedurally, which requires the imaging core of a catheter to be pulled back at a known uniform speed,
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- two-dimensions, longitudinal reconstructions of the anatomy is desired.
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### **Intended Use/Indications for Use**

The IVUS modality of the iLab™ Polaris Multi-Modality Guidance System is intended for ultrasound examinations of intravascular pathology. Intravascular ultrasound is indicated in patients who are candidates for transluminal interventional procedures such as angioplasty and atherectomy.

FFR and DFR™ are intended for use in catheterization and related cardiovascular specialty laboratories to compute, and display various physiological parameters based on the output from one or more electrodes, transducers, or measuring devices.

FFR and DFR are indicated to provide hemodynamic information for use in the diagnosis and treatment of patients that undergo measurement of physiological parameters.

The Imaging Catheters generate ultrasound images and are intended for ultrasound examination of vascular and cardiac pathology. Boston Scientific manufactures a wide variety of catheters for different applications. The recommended use of each of these catheters may vary depending on the size and type of the catheter. Please refer to the Imaging Catheter Directions for Use, packaged with each catheter.

#### **Indications for Auto Pullback Use (IVUS Only)**

Automatic Pullback is indicated when the following occurs:

- The physician/operator wants to standardize the method in which intravascular ultrasound images are obtained and documented: procedure-to-procedure, operator-to-operator.
  - The physician/operator wants to make linear distance determinations post-procedurally, which requires the imaging core of a catheter to be pulled back at a known uniform speed.
  - Two-dimensional, longitudinal reconstruction of the anatomy is desired.
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## Comparison of Technological Characteristics

The iLab Polaris Multi-Modality Guidance System with updated MDU5+ Motor Drive Unit supporting additional Auto Pullback functionality is substantially equivalent to the predicate iLab Polaris Multi-Modality Guidance System (K201178) in intended use, fundamental design technology, functional modalities and performance features. Additional auto pullback options for user selection do not raise different questions of safety or effectiveness.

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## Non-clinical Performance

Determination of substantial equivalence is based on an assessment of non-clinical performance data which includes hardware verification and validation carried out using the updated MDU5+ Motor Drive Unit. Testing was conducted according to applicable international standards, FDA recognized consensus standards, and the same well-established test methods and criteria applied to the predicate device. The following standards and guidance are applicable in demonstration of substantial equivalence related to the testing:

- EN 62304:2006+AMD 1:2015-/IEC 62304 (Edition 1.1 2015-06) (FDA/CDRH Recognition No.-13-79) Medical Device Software - Software Life-Cycle Processes
- ANSI AAMI BP-22:1994 (R) 2011 (FDA/CDRH Recognition No.3-44) - Blood Pressure Transducers
- ANSI AAMI ES 60601-1: 2005/(R)2012 and A1:2012 (FDA/CDRH Recognition No.19-4) - Medical Electrical Equipment- Part1: General requirements for Basic Safety and Essential Performance
- IEC 60601-1-2:2014 4th Ed. (FDA/CDRH Recognition No.19-8) Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests

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## Clinical Performance Data

Not applicable. A determination of Substantial Equivalence for this modification is not based on clinical data. Substantial Equivalence is based on non-clinical performance data.

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## Conclusion

Based on a comparison of intended use, fundamental design technology, functional modalities and performance features the iLab Polaris Multi-Modality Guidance System with updated MDU5+ Motor Drive Unit including additional Auto Pullback speeds is substantially equivalent to the predicate device. A comparison of the modified and predicate devices, along with verification and validation testing applicable to the modified device, supports a conclusion of substantial equivalence and raise no new issues of safety and effectiveness.

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