



December 8, 2022

InkSpace Imaging, Inc.
% Taras Bouzakine
Director
Veranex, Inc.
224 Airport Parkway, Suite 250
SAN JOSE CA 95110

Re: K223487
Trade/Device Name: InkSpace Imaging Body Array
Regulation Number: 21 CFR 892.1000
Regulation Name: Magnetic resonance diagnostic device
Regulatory Class: Class II
Product Code: MOS
Dated: November 18, 2022
Received: November 21, 2022

Dear Taras Bouzakine:

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) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

A handwritten signature in black ink, appearing to read 'D. Krainak', is written over a large, light blue watermark of the letters 'FDA'.

Daniel M. Krainak, Ph.D.
Assistant Director
Magnetic Resonance and Nuclear Medicine Team
DHT8C: Division of Radiological Imaging
and Radiation Therapy Devices
OHT8: Office of Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)

K223487

Device Name

InkSpace Imaging Body Array

Indications for Use (Describe)

Pediatric Population:

The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of pediatric cardiac and body in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.

Adult Population:

The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of general human anatomy in adults, such as spine, shoulder, elbow, knee, foot, and prostate in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.

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.

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510(k) SUMMARY

510(k) Notification K 223487

GENERAL INFORMATION [807.92(a)(1)]

Applicant:

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Contact Person:

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Date Prepared: November 18, 2022

DEVICE INFORMATION [807.92(A)(2)]

Classification:

Class II per 21 CFR§892.1000

Product Code:

MOS

Trade Name:

InkSpace Imaging Body Array

Generic/Common Name:

Coil, Magnetic Resonance, Specialty

PREDICATE DEVICE(S) [807.92(A)(3)]

InkSpace Imaging Pediatric Body Array (K213397)

DEVICE DESCRIPTION [807.92(A)(4)]

The InkSpace Imaging Body Array is a phased array, receive-only coil intended to work with GE 3.0T MRI scanners to provide optimal signal-to-noise ratio and uniformity for high resolution body MR imaging examinations. The size and layout of individual channels of the InkSpace Imaging Body Array were designed to fit the pediatric and adult populations for body imaging. The coil is comprised of 24 total channels, with two individual flexible pads, intended to be

510(k) SUMMARY

placed anterior and posterior to the patient, containing 12 individual elements. The elements are optimized to be flexible to conform to pediatric and adult patients’ anatomies.

INDICATIONS FOR USE [807.92(a)(5)]

Pediatric Population:

The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of pediatric cardiac and body in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.

Adult Population:

The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of general human anatomy in adults, such as spine, shoulder, elbow, knee, foot, and prostate in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.

COMPARISON OF TECHNOLOGICAL CHARACTERISTICS WITH THE PREDICATE DEVICES [807.92(a)(6)]

The technological characteristics of the modified InkSpace Imaging Body Array are substantially equivalent to the predicate device, InkSpace Imaging Pediatric Body Array (K213397). Table 1 lists the technological characteristics of the predicate and modified devices and provides rationale to support a determination of substantial equivalence. Any differences between the devices do not raise any new or different questions of safety or effectiveness.

Table 1: Summary of Technological Characteristics

Characteristics	InkSpace Imaging Pediatric Body Array (K213397)	InkSpace Imaging Body Array	Substantial Equivalence Rationale
Classification	Magnetic Resonance Diagnostic Device per 21 CFR§892.1000	Magnetic Resonance Diagnostic Device per 21 CFR§892.1000	N/A (Same).
Product Code	MOS (Coil, Magnetic Resonance, Specialty)	MOS (Coil, Magnetic Resonance, Specialty)	N/A (Same).
Intended Use	The InkSpace Imaging Pediatric Body Array is a receive-only RF coil designed for use with GE 3.0T MRI systems. It is used for obtaining diagnostic images of the body and cardiac regions of pediatric patients in magnetic resonance imaging systems. The nucleus detected is hydrogen. Anatomic Regions: Cardiac/Body for pediatric patients.	<p>Pediatric Population: The InkSpace Imaging Body Array is a receive-only RF coil designed for use with GE 3.0T MRI systems. It is used for obtaining diagnostic images of the body and cardiac regions of pediatric patients in magnetic resonance imaging systems. The nucleus detected is hydrogen. Anatomic Regions: Cardiac/Body for pediatric patients.</p> <p>Adult Population: The InkSpace Imaging Body Array is a receive-only RF coil designed for use with GE 3.0T MRI systems. It is used for obtaining diagnostic images of general human anatomy such as spine, shoulder, elbow, knee, foot, and prostate for adult</p>	Intended use of the modified device and predicate device maintain the same general purpose and function.

Table 1: Summary of Technological Characteristics (Cont.)

Characteristics	InkSpace Imaging Pediatric Body Array (K213397)	InkSpace Imaging Body Array	Substantial Equivalence Rationale
		patients in magnetic resonance imaging systems. The nucleus detected is hydrogen.	
Indications for Use	The InkSpace Imaging Pediatric Body Array is a receive-only coil, used for obtaining diagnostic images of pediatric cardiac and body in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.	<p>Pediatric Population: The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of pediatric cardiac and body in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.</p> <p>Adult Population: The InkSpace Imaging Body Array is a receive-only coil, used for obtaining diagnostic images of general human anatomy in adults, such as spine, shoulder, elbow, knee, foot, and prostate in GE 3.0T magnetic resonance imaging systems. These images, when interpreted by a trained physician, yield information that may assist in diagnosis.</p>	Patient population updated from pediatric only to pediatric and adult.
Patient Anatomy and Population	Imaging of general body part regions including cardiac regions; designed for pediatric patients.	Imaging of general body part regions including cardiac regions; intended for pediatric patients. Imaging of general human anatomy, such as spine, shoulder, elbow, knee, foot, and prostate; intended for adult patients.	Patient population updated from pediatric only to pediatric and adult.
Comparison of Technological Characteristics	24 channel, receive-only phased array, designed for 3.0T MR systems (127.73 MHz, hydrogen). Device is reusable, non-sterile, and by prescription only. Device is made of soft, pliable materials, and is composed of an anterior and a posterior pad.	24 channel, receive-only phased array, designed for 3.0T MR systems (127.73 MHz, hydrogen). Device is reusable, non-sterile, and by prescription only. Device is made of soft, pliable materials, and is composed of an anterior and a posterior pad.	N/A (Same).
Compatible MRI Systems	GE 3.0T MRI Systems	GE 3.0T MRI Systems	N/A (Same).
Biocompatibility	ISO 10993 testing	ISO 10993 testing	N/A (Same).

SUBSTANTIAL EQUIVALENCE

The InkSpace Imaging Body Array is substantially equivalent to the predicate device, InkSpace Imaging Pediatric Body Array with regard to function, intended use, and physical characteristics. There are no differences in the technological characteristics between the devices which would raise any different questions of safety or efficacy. Thus, the modified InkSpace Imaging Body Array is substantially equivalent to the predicate device.

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PERFORMANCE DATA [807.92(b)]

Performance testing was performed in accordance with the FDA Guidance Document entitled, “*Magnetic Resonance (MR) Receive-only Coil – Performance Criteria for Safety and Performance Based Pathway*,” issued December 11, 2020.

All applicable nonclinical testing conducted on the InkSpace Imaging Pediatric Body Array (predicate device) is still applicable to the modified InkSpace Imaging Body Array.

Additional analysis of acquired image quality was conducted based on sample clinical images from adult patients. These images, as well as a review of their clinical quality by a board-certified radiologist, have been included as part of this submission.

[807.92(b)(1)] Nonclinical Testing Summary:

The nonclinical, bench testing previously performed on the InkSpace Imaging Pediatric Body Array:

- Image Signal to Noise (SNR)
- Image Uniformity
- Surface Heating
- Decoupling Circuit
- EMC – Immunity, Electrostatic Discharge
- General Electrical/Mechanical Safety

Additional testing performed on the InkSpace Imaging Body Array:

- Acquired Image Quality

In addition to the above nonclinical testing, the InkSpace Imaging Pediatric Body Array also previously underwent the following testing:

- Biocompatibility Testing (Cytotoxicity, Sensitization, Irritation)
- Usability Testing
- Transit Testing

The collective results of the nonclinical testing demonstrate that the materials chosen, the manufacturing processes, and design of the InkSpace Imaging Body Array meet the established specifications necessary for consistent performance during its intended use. In addition, the collective bench testing demonstrates that the InkSpace Imaging Body Array does not raise different questions of safety or effectiveness for MR imaging examinations when compared to the predicate device.

[807.92(b)(2)] Clinical Testing Summary:

Not applicable. Clinical testing was not required to demonstrate substantial equivalence of InkSpace Imaging Body Array to the predicate device. Acquired image quality from adult patients was analyzed based on sample clinical images. These images, as well as a review of their clinical quality by a board-certified radiologist, have been included as part of this submission.

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CONCLUSIONS [807.92(b)(3)]

Based on the results from the nonclinical tests performed in support of the InkSpace Imaging Body Array, it is concluded that the modified device is safe, effective, and performs at least as safely and effectively as the legally marketed predicate device.

SUMMARY

The InkSpace Imaging Body Array is substantially equivalent to the predicate device.