

iRay Technology Taicang Ltd.
% Mr. Junjie Qian
Registration & Regulation Affairs Engineer
No. 33 Xinggang Rd., Taicang Port Economic &
Technological Development Zone
Taicang, Jiangsu 215434
CHINA

May 12, 2020

Re: K201043

Trade/Device Name: Wireless Digital Flat Panel Detector (Model: Mars1717V-VSI, Mano4343W)

Regulation Number: 21 CFR 892.1680 Regulation Name: Stationary x-ray system

Regulatory Class: Class II Product Code: MQB Dated: April 8, 2020 Received: April 20, 2020

Dear Mr. Qian:

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 <u>Federal Register</u>.

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-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">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,

For

Thalia T. Mills, Ph.D.
Director
Division of Radiological Health
OHT7: Office of In Vitro Diagnostics
and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

510(k) Number (if known)

Form Approved: OMB No. 0910-0120

Expiration Date: 06/30/2020 See PRA Statement below.

K201043	
Device Name Wireless Digital Flat Panel Detector (Model:Mars1717V-VSI, Mano43-	43W)
Indications for Use (Describe) Wireless Digital Flat Panel Detector (Model:Mars1717V-VSI, M designed to provide general radiographic diagnosis for human an are intended to replace film/screen systems in all general–purposintended for mammography, dental applications.	atomy including both adult and pediatric patients. They
There are no differences between the 2 models except the model	name and label.
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 SEPARAT	E PAGE IF NEEDED.

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510 (k) SUMMARY OF SAFETY AND EFFECTIVENESS

K201043

510 (k) SUMMARY OF SAFETY AND EFFECTIVENESS

(As Required by 21 CFR 807.92)

1. <u>Date Prepared [21 CFR 807.92(a)(1)]</u>

April 8th, 2020

2. Submitter's Information [21 CFR 807.92(a)(1)]

<u>Company</u> Name: iRay Technology Taicang Ltd.

Company Address: No.33 Xinggang Road, Taicang Port Economic and

Technological Development Zone, Jiangsu, China 215434

Contact Person: Junjie Qian

Phone: 0512-53690872 **Fax:** 0512-53690872

Email: junjie.qian@iraygroup.com

3. Trade Name, Common Name, Classification [21 CFR 807.92(a)(2)]

<u>Trade Name:</u> Wireless Digital Flat Panel Detector

Common Name: Solid State X-Ray Imager

Model Name: Mars1717V-VSI

Mano4343W

Classification Name: Stationary x-ray system

Product Code: MQB

Regulation Number: 21 CFR 892.1680

Device Class: Class II

4. Identification of Predicate Devices(s) [21 CFR 807.92(a)(3)]

The identification predicates within this submission are as follows:

Manufacturer: iRay Technology Taicang Ltd.

Trade Name: Wireless Digital Flat Panel Detector

Model Name: Mars1717XF-CSI

Product Code: MQB

Device Class: Class II

Classification Name: Stationary x-ray system

FDA 510 (k) #: K183713

5. Description of the Device [21 CFR 807.92(a)(4)]

Mars1717V-VSI and Mano4343W Wireless Digital Panel Detectors (Hereinafter referred to as Mars1717V-VSI and Mano4343W) are the kind of wireless digital flat panel detectors. They support the single frame mode, with the key component of TFT/PD image sensor flat panel of active area: 427mm x 427mm. Two models Mars1717V-VSI and Mano4343W are totally same except for label and model name. The sensor plate of Mars1717V-VSI and Mano4343W is direct-deposited with CsI scintillator to achieve the conversion from X-ray to visible photon. The visible photons are transformed to electron signals by diode capacitor array within TFT panel, which are composed and processed by connecting to scanning and readout electronics, consequently to form a panel image by transmitting to PC through the user interface. The major function of the Mars1717V-VSI and Mano4343W is to convert the X-ray to digital image, with the application of high-resolution X-ray imaging. Both kinds of detectors are the key component of DR system, enable to complete the digitalization of the medical X-ray imaging with the DR system software.

iRay SDK(include iDetector) is intend to supply API interface for DR system manufacturers.DR system manufacturer control the detector by SDK interface. SDK is not intended to be used directly by other users beside DR system manufacturers. The iRay SDK is unchanged from the predicate device.

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6. <u>Intended Use [21 CFR 807.92(a)(5)]</u>

6.1. Indications for use

Mars1717V-VSI and Mano4343W are indicated for digital imaging solutions designed to provide general radiographic diagnosis for human anatomy including both adult and pediatric patients. They are intended to replace film/screen systems in all general—purpose diagnostic procedures. These two devices are not intended for mammography, dental applications. There are no differences between the 2 models except the model name and label.

6.2. Suitable patient

They are suitable for providing digital X-ray imaging for DR system to provide general radiographic diagnosis for human anatomy including both adult and pediatric patients, but not intended for mammography or dental applications. The remaining notes depend on the DR system.

6.3. Processing of input and output

When flat panel detector works continuously, it can automatically distinguish X-ray and output an imaging for diagnosis of disease, injury, or of any applicable health problem.

7. Technological Characteristic [21 CFR 807.92(a)(6)]

Item	Predicate Device: Wireless Digital Flat Panel Detector	Proposed Device: Wireless Digital Flat Panel Detector
510(K) Number	K183713	K201043
Intended Use	The Mars1717XF-CSI Wireless Digital Flat Panel Detector is indicated for digital imaging	same

	Predicate Device:	Proposed Device:
Item	Wireless Digital Flat Panel	Wireless Digital Flat Panel
	Detector	Detector
	solution designed for providing	
	general radiographic system in all	
	general-purpose diagnostic	
	procedures.	
	Mars1717XF-CSI Wireless	Same with Mars1717XF-CSI,
	Digital Flat Panel Detector is	additionally, in the
	indicated for digital imaging	consideration of patient size
	solution designed for providing	(i.e., height, weight, body part
	general radiographic diagnosis of	thickness) and usable dose
	human anatomy.	range, the detector could be
		used for general X ray
	It is intended to replace	diagnosis of usual body part
	radiographic film/screen systems	for adults and pediatric
	in all general-purpose diagnostic	patients.
Indications for	procedures. This device is not	
Use	intended for mammography or	
	dental applications.	
	We understand the Agency has	
	become aware of situations where	
	solid state detectors inserted into	
	radiographic systems adversely	
	impacted device performance due	
	to improper integration	
	(reference:http://www.fda.gov/do	
	wnloads/MedicalDevices/Resourc	

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	Predicate Device:	Proposed Device:
Item	Wireless Digital Flat Panel	Wireless Digital Flat Panel
	Detector	Detector
	esforYou/Industry/UCM385149.p	
	df).	
	Below is a summary of the	
	information from the	
	Mars1717XF-CSI user manuals	
	covering key electromechanical	
	and computer requirements	
	needed for X-ray system interface	
	and integration.	
	1. Mechanical interface	
	requirements.	
	2. Computer requirements	
	3. Data communication interface	
	requirements	
	4. Electrical power requirements	
	5. X-ray trigger interface	
	requirements	
	Neither the Mars1717XF-CSI	
	detector nor its software act as an	
	X-ray generator controller, and	
	therefore, the device is not subject	
	to Electronic Product Radiation	
	Control (EPRC) performance	
	standards and reporting	
	requirements.	

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	Predicate Device:	Proposed Device:
Item	Wireless Digital Flat Panel	Wireless Digital Flat Panel
	Detector	Detector
Classification	Stationary x-ray system	Same
Name	Stationary X-ray system	Same
Product Code	MQB	Same
Regulation	21 CFR 892.1680	Same
Number	21 CI K 0/2.1000	Same
Panel:	Radiology	Same
Classification:	II	Same
X-Ray Absorber	CsI	Same
(Scintillator):		Same
Installation	Wireless, Portable	Same
Type:	Wholess, Fordore	
Readout	Thin Film Transistor	Same
Mechanism:		
Image Matrix	2832×2836 pixels	3072 ×3072 pixels
Size:	2002 ·· 2000 pinois	3072 ×3072 pixels
Pixel Pitch:	150μm	139µm
ADC	16 bit	Same
Digitization		
Effective	424.8 mm ×425.4 mm	427 mm ×427 mm
Imaging Area:	727.0 mm × 723.7 mm	127
Spatial	3.3lp/mm	3.6lp/mm
Resolution:	1	I.
Modulation		
Transfer	0.49 at 1 lp/mm	0.65 at 1 lp/mm
Function	0.17 tt 1 1p/111111	order i iprimi
(MTF)		

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	Predicate Device:	Proposed Device:
Item	Wireless Digital Flat Panel	Wireless Digital Flat Panel
	Detector	Detector
Detective		
Quantum	0.40 at 1.1n/mm (POA5, 2.5u/Cy)	Same
Efficiency	0.40 at 1 lp/mm (RQA5, 2.5μGy)	Same
(DQE)		
Power	Max. 20W	Max. 30W
Consumption:	WIGA. 20 W	Wax. 50 W
		a) Wired (only for
Communication		service): Gigabit
s:	Wireless: IEEE 802.11a/b/g/n (2.4	Ethernet (1000BASE-T)
(Wireless	GHz / 5 GHz)	b) Wireless: IEEE
functionality)		802.11a/b/g/n/ac (2.4 GHz
		/ 5 GHz)
Imaging protect Plate:	Carbon Fiber Plate	Same
Cooling:	Air cooling	Same
Dimensions:	460 mm ×460 mm ×15 mm	460 mm ×460 mm ×15.3 mm
	Temperature: +5 ~ +30°C	Temperature: +5 ~ +35 °C
	Humidity: 10 ~ 80%	Humidity: 5 ~ 95%
Operation:	(Non-Condensing)	(Non-Condensing)
	Atmospheric pressure: 70 ~ 106	Atmospheric pressure: 55 ~
	kPa	106 kPa
	Altitude: Max. 3000 meters	Altitude: Max. 3000 meters
Storage and	Temperature: -20 ~ +50 °C	Temperature: -10 ~ +55 °C
Transportation:	Humidity: 10 ~ 90%	Humidity: 5 ~ 95%

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	Predicate Device:	Proposed Device:
Item	Wireless Digital Flat Panel	Wireless Digital Flat Panel
	Detector	Detector
(detector)	(Non-Condensing)	(Non-Condensing)
	Atmospheric pressure: 70 ~ 106	Atmospheric pressure: 55 ~
	kPa	106 kPa
	Altitude: Max. 3000 meters	Altitude: Max. 3000 meters
	iRay SDK(include iDetector) is	
	intend to supply API interface for	
	DR system manufacturers. DR	
Software	system manufacturer control the	Same
Software	detector by SDK interface. SDK	Same
	is not intended to use directly by	
	other users beside DR system	
	manufacturers.	
	1. Guidance for the Submission	
	of 510(k)'s for Solid State X-	
	ray Imaging Devices;	Come with Marc 1717VE CCI
	2. The 510(k) Program:	Same with Mars1717XF-CSI,
	Evaluating Substantial	additionally:
Utilized FDA	Equivalence in Premarket	1. Guidance for "Premarket
guidance	Notifications[510(k)];	Assessment of Pediatric
documents	3. Content of Premarket	Medical Devices";
	Submissions for Management	2. Guidance for "Pediatric
	of Cybersecurity in Medical	Information for X-ray
	Devices;	Imaging Device Premarket
	4. Radio Frequency Wireless	Notifications".
	Technology in Medical	
	Devices.	

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8. System requirements to operate with other radiographic system components

1) Recommended Generator Specification:

Energy range: 40~150keV

mA range: 10~1000mA (depending on the generator power)

ms range: 10~6300ms to produce 0.1~1000mAs (depending on the generator power) Note: To our best knowledge, the detector is compatible with the X-ray generators with the specifications described above. If you have any questions regarding the

compatibility issue for other generators, please contact your distributor or iRay's

service office.

Application Program Interface (API) for system integration manufacturer
 Peripheral hardware: Mars1717V-VSI and Mano4343W are connected via wireless communication.

recommended computer hardware requirement is:

- Windows 7 32/64 bit
- •Intel Core i7 3.6G
- •4G DDR3
- •640 G
- Intel Pro EXP9301CT PRO, Gigabit Network Adapter with PCIe interface
- 3) X-ray exposure mode

The AED trigger module is a unit can connect X-ray signal in the Mars1717V-VSI and Mano4343W. Once there is X-ray generator exposure exist, the AED trigger module will detect the X-ray radiation and output signal to the detector. Until the exposure finished, the detector will receive a signal which represent the end of exposure from the inner trigger module and begin to acquire the image.

9. Substantial Equivalence [21 CFR 807.92(b)(1) and 807.92(b)(2)]

1) Electrical Safety and EMC testing:

Electrical, mechanical, environmental safety test according to IEC/ES 60601-1 was performed, and EMC testing was also conducted in accordance with IEC 60601-1-2. All test results are meet the standard requirements.

2) Biological Evaluation:

The materials of the detector which contact operators' or patient's skin have been evaluated with the ISO 10993-1. And the evaluation results and test result assured the safety the same as the predicate device.

3) Non-clinical Considerations:

One modification from the predicate device to Mars1717V-VSI and Mano4343W is geometric design, related to Amorphous Silicon (A-Si) panel size and structure size design. Another modification is wireless functionality, proposed device has higher wireless transmission speed.

The non-clinical studies have been performed and the results have shown that sections of the non-clinical consideration mentioned in the 'Guidance for the Submission of 510(k)s for Solid State X-ray Imaging Devices' are substantially equivalent to the non-consideration of predicate devices on the Market (Mars1717XF-CSI, K183713).

4) Clinical Consideration:

Intended use, fundamental scientific technology, regulatory requirement, non-clinical performance, labeling, quality-assurance program and software keep the same with those of predicate device. Additionally, as mentioned in clinical considerations in 'Guidance for the Submission of 510(k)s for Solid State X-ray Imaging Devices', clinical consideration may not necessary for changes in the pixel size and resolution with the same x-ray detection material and may not necessary for changes in the wireless functionality if non-clinical information is sufficient to support the substantial equivalence.

There was no significant difference between the images of the Mars1717V-VSI/Mano4343W and those of the predicate device.

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10. Conclusion [21 CFR 807.92(b)(3)]

In accordance with the Federal Food, Drug and Cosmetic Act, 21 CFR Part 807 and based on the information provided in this premarket notification, iRay Technology Taicang Ltd. concludes that Mars1717V-VSI and Mano4343W are substantially equivalent to predicate device with regards to safety and effectiveness.