July 13, 2022



LG Electronics Inc. % Woo Chai Kyoung Senior Consultant GMS Consulting 4th Floor, Digital Cube, 34, Sangamsan-ro Seoul, Mapo-gu 03909 KOREA

Re: K221434

Trade/Device Name: 17HQ701G-B Regulation Number: 21 CFR 892.1680 Regulation Name: Stationary x-ray system Regulatory Class: Class II Product Code: MQB Dated: May 16, 2022 Received: May 17, 2022

Dear Woo Chai Kyoung:

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 <a href="https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm">https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm</a> 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 <a href="https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products">https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products</a>); 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 <u>https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems</u>.

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

Sincerely,

Laurel Burk, Ph.D. Assistant Director Diagnostic X-Ray Systems Team DHT8B: Division of Radiological Imaging Devices and Electronic Products 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)* K221434

Device Name 17HQ701G-B

Indications for Use (Describe)

The Flat Panel Digital X-ray Detector 17HQ701G-B is indicated for digital imaging solution designed for general radiographic system for human anatomy. It is intended to replace film or screen based radiographic systems in all general purpose diagnostic procedures. Not to be used for mammography.

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

[As Required by 21 CFR 807.92]

### 1. Date Prepared [21 CFR 807.92(a)(a)]

May 16, 2022

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

•	Name of Sponsor: - Address:	LG Electronics Inc. 77, Sanho-daero, Gumi-si, Gyeongsangbuk-do, 39381, Republic of Korea
•	Name of Manufacturer: - Address:	LG Electronics Inc. 77, Sanho-daero, Gumi-si, Gyeongsangbuk-do, 39381, Republic of Korea
•	Contact Name: - Telephone No.: - Email Address:	Jinhwan Jun / Chief Research Engineer +82-31-8066-5641 jinhwan.jun@lge.com

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

Trade/Device/Model Name	17HQ701G-B
Common Name	Flat Panel Digital X-ray Detector
Device Classification Name Stationary X-ray System	
Regulation Number	21 CFR 892.1680
Classification Product Code	MQB
Device Class	П
510(k) Review Panel	Radiology

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

The identified predicate devices within this submission are shown as follow;

#### Predicate Device

•	510(k) Number:	K182348
•	Applicant:	LG Electronics Inc.
•	Trade/Device Name:	14HK701G-W
•	Common Name:	Flat Panel Digital X-ray Detector
•	Classification Name:	Stationary X-ray System
•	Regulation Number:	21 CFR 892.1680
•	Classification Product Code	MQB
•	Device Class:	II
•	510(k) Review Panel:	Radiology

The predicate devices have not been subject to a design-related recall

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

This model is an x-ray imaging device, a system that can acquire and process X-ray images as digital images. It utilizes amorphous silicon and a high-performance scintillator to ensure sharp high-definition image quality with the resolution of 3.6 lp/mm and the pixel pitches of 140 um. This device is a flat panel based X-ray image acquisition device. This device must be used in conjunction with an operating PC and an X-ray generator. This device can be used for digitizing and transferring X-ray images for radiological diagnosis. The data transmission between the Detector and PC can be enabled with a wired (cable) or wireless connection.

## 6. Indications for use [21 CFR 807.92(a)(5)]

The Flat Panel Digital X-ray Detector 17HQ701G-B is indicated for digital imaging solution designed for general radiographic system for human anatomy. It is intended to replace film or screen based radiographic systems in all general purpose diagnostic procedures. Not to be used for mammography.

### 7. Intended Use [21 CFR 807.92(a)(5)]

The Flat Panel Digital X-ray Detector 17HQ701G-B is a prescription device, and it is not intended to be used for mammography.

- The detector is indicated for digital imaging solution designed for general radiographic system for human anatomy
- The detector is indicated to replace film or screen based radiographic systems in all general purpose diagnostic procedures.

## 8. Technological Characteristics (Equivalence to Predicate Device) [21 CFR 807.92(a)(6)]

There are no significant differences in the technological characteristics of these devices compared to the predicate devices which adversely affect safety or effectiveness. Provided below is a table summarizing and comparing the technological characteristics of the 17HQ701G-B and the predicate devices:

	Proposed Device	Predicate Device	Note
K Number	TBD	K182348	-
Manufacturer	LG Electronics Inc.	LG Electronics Inc.	Same
Trade Name	17HQ701G-B	14HK701G-W	-
Common Name	Flat Panel Digital X-ray Detector	Flat Panel Digital X-ray Detector	Same
Product Code	MQB	MQB	Same
Regulation	21 CFR 892.1680	21 CFR 892.1680	Same
Number			
510(k) Review	Radiology Radiology S		Same
Panel			
Indications for	The Flat Panel Digital X-ray Detector	Flat Panel Digital X-ray Detectors are	Same
Jse 17HQ701G-B is indicated for digital indicated for digital imaging solution			
	imaging solution designed for general	designed for general radiographic	
	radiographic system for human	system for human anatomy. It is	
	anatomy. It is intended to replace	intended to replace film or screen	
	film or screen based radiographic	based radiographic systems in	
	systems in all general purpose	general purpose diagnostic	
	diagnostic procedures. Not to be	procedures all and not to be used for	
	used for mammography.	mammography.	
Intended use	The Flat Panel Digital X-ray Detector	The Flat Panel Digital X-ray Detector	Same
	17HQ701G-B is a prescription device,	14HK701G-W is a prescription device,	
	and it is not intended to be used for	and it is not intended to be used for	
	mammography.	mammography.	
	<ul> <li>The detector is indicated for</li> </ul>	<ul> <li>The detector is indicated for</li> </ul>	
	digital imaging solution	digital imaging solution	
	designed for general	designed for general	
	radiographic system for	radiographic system for	
	human anatomy	human anatomy	
	<ul> <li>The detector is indicated to</li> </ul>	<ul> <li>The detector is indicated to</li> </ul>	
	replace film or screen based	replace film or screen based	
	radiographic systems in all	radiographic systems in all	
	general purpose diagnostic	general purpose diagnostic	
	procedures.	procedures.	
Detector		1	
Scintillator	CsI	CsI	Same
Imaging Area	17 x 17 inches	13.7 x 16.8 inches	Different
Pixel Matrix	3,072 x 3,072 pixels	2,500 x 3,052 pixels	Different
Pixel Pitch	140 um	140 um	Same
High Contrast	3.6 lp/mm	3.6 lp/mm	Same
Limiting			
Resolution			
(LP/mm)			
Communication	Wired/Wireless	Wired/Wireless	Same
DQE	Typ.66% @0.1lp/mm	Typ.72% @0.1lp/mm	Different
MTF	Typ.84% @0.5lp/mm	Typ.89% @0.5lp/mm	Different

	Proposed Device	Predicate Device	Note
Resolution	on 3.6lp 3.6lp		Same
Anatomical Sites	General	General	Same
Exposure Mode	Manual, Auto (AED)	Manual, Auto (AED)	Same
Semi Dynamic mode	,		Same
Wireless	Standard:Standard:802.11 a/b/g/n/ac compliance802.11 a/b/g/n/ac complianceFrequency: 2.4 GHz/5GHzFrequency: 2.4 GHz/5GHzBandwidth: 20MHz/40MHz/80MHzBandwidth: 20MHz/40MHz/80MHzMIMO: 2x2MIMO: 2x2		Same
Rating	ng 24V 2.1A 24V 2.1A Sa		Same
Gap Analysis	Cap Analysis There is some difference in the 'Image Area', 'Pixel Matrix', 'DQE' and 'MTF'. Image Area, Pixel Matrix, DQE and MTF are not related to the device's 'safety and 'performance.' So Proposed device (17HQ701G-B) and Predicate Device (14HK701G-W) are substantially same.		

\*\* Semi Dynamic mode is a function that can transmit five images per second to the PC.

There are no significant differences between the 17HQ701G-B and the predicate device that would adversely affect the use of the product. It is substantially equivalent to these devices in design, function, materials, operational principles and intended use. The proposed device, 17HQ701G-B has been tested about electrical safety, EMC and performance, and the software has been validated. In addition, the clinical data has been provided to support the substantial equivalence to the predicate devices.

#### 9. Non-Clinical Test summary

The 17HQ701G-B comply with voluntary standards for electrical safety, electromagnetic compatibility. The following data were provided in support of the substantial equivalence determination:

1) Electrical Safety, Electromagnetic Compatibility and Performance:

The 17HQ701G-B comply with the electrical safety and electromagnetic compatibility requirements established by the standards.

Standards No.	Standards Organization	Standard Title	Version	Publication Year
ES60601-1	AAMI	Medical Electrical Equipment - Part 1: General Requirements For Basic Safety And Essential Performance (IEC 60601-1:2005, MOD)	ES60601-1: 2005(R)201 2 and A1:2012	2014
60601-1-2	IEC	Medical Electrical Equipment - Part 1-2: General Requirements for Safety – Collateral Standard: Electromagnetic Compatibility - Requirements and Tests	nts for Safety – Collateral Standard:	
-	FDA	Radio Frequency Wireless Technology in Medical Devices	August 14	2013

#### 2) Software Validation

The 17HQ701G-B contains MODERATE level of concern software as firmware. The software was designed and developed according to a software development process and was verified and validated. Software information is provided in accordance with FDA guidance:

• The content of premarket submissions for software contained in medical devices, on May 11, 2005

#### 3) Biocompatibility

• ISO 10993-1 and series, Biological evaluation of medical devices

#### 4) Performance Test

Imaging performance test has been conducted according to:

• IEC 62220-1, Medical Electrical Equipment – Characteristics of Digital X-ray Imaging Devices – Part 1-1: Determination of the Detective Quantum Efficiency – Detectors Used in Radiographic Imaging.

#### 5) Cybersecurity

- Content of Premarket Submissions for Management of Cybersecurity in Medical Devices, on October 18, 2018
- Postmarket Management of Cybersecurity in Medical Devices, on December 28, 2016

- 6) Label
- CFR Part 801
- Pediatric Information for X-ray Imaging Device Premarket Notifications, on November 28, 2017

#### **10. Clinical Test Summary**

Clinical data has been provided according to FDA guidance document "Guidance for the Submission of 510(k)s for Solid Sate X-ray Imaging Devices". The data was not necessary to establish substantial equivalence based on the modifications to the device but provided further evidence in addition to the laboratory performance data to show that the device works as intended.

### 11. Substantial Equivalence [21 CFR 807.92(b)(1) and 807.92]

There are no significant differences between 17HQ701G-B and the predicate device, K182348 that would adversely affect the use of the product. It is substantially equivalent to these devices in indications for use and technology characteristics.

### 12. Conclusion [21 CFR 807.92(b)(3)]

In according with the Federal Food & Drug and cosmetic Act, 21 CFR Part 807, and based on the information provided in this premarket notification LG Electronics, concludes that the 17HQ701G-B is substantially equivalent in safety and effectiveness to the predicate device as described herein.