



Food and Drug Administration  
10903 New Hampshire Avenue  
Document Control Center – WO66-G609  
Silver Spring, MD 20993-0002

Samsung Electronics Co., Ltd.  
% Chulsin Kim  
Regulatory Affairs Manager  
129, Samsung-ro, Yeongtong-gu  
Suwon-si, Gyeonggi-do 443-742  
REPUBLIC OF KOREA

April 2, 2015

Re: K150165  
Trade/Device Name: GC85A  
Regulation Number: 21 CFR 892.1680  
Regulation Name: Stationary x-ray system  
Regulatory Class: II  
Product Code: KPR  
Dated: January 23, 2015  
Received: January 26, 2015

Dear Chulsin Kim:

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. 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); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638 2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>. 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 the CDRH’s Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,

A handwritten signature in black ink that reads "Robert A. Ochs". The signature is written in a cursive style. Behind the signature, there is a faint, semi-transparent watermark of the FDA logo.

Robert Ochs, Ph.D.  
Acting 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)

**K150165**

Device Name

GC85A

Indications for Use (Describe)

The GC85A Digital X-ray Imaging System is intended for use in generating radiographic images of human anatomy by a qualified/trained doctor or technician. This device is not intended for mammographic applications.

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.

**\*DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.\***

The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services  
Food and Drug Administration  
Office of Chief Information Officer  
Paperwork Reduction Act (PRA) Staff  
[PRASStaff@fda.hhs.gov](mailto:PRASStaff@fda.hhs.gov)

*"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."*



## Section 5: 510(k) Summary

This summary of 510(k) safety and effectiveness information is being submitted accordance with requirements of 21 CFR 807.92

1. **Date:** January 23, 2015
2. **Submitter**
  - A. Company Name: SAMSUNG ELECTRONICS Co., Ltd.
  - B. Address: 129, Samsung-ro, Yeongtong-gu, Suwon-si, Gyeonggi-do, 443-742, Republic of Korea
3. **Primary Contact Person**
  - A. Name: KIM, CHULSIN
  - B. Title: Regulatory Affairs Manager
  - C. Phone Number: +82-31-200-7661
  - D. FAX Number: +82-31-200-1199
  - E-Mail: [chulsin.kim@samsung.com](mailto:chulsin.kim@samsung.com)
4. **Secondary Contact Person**
  - A. Name: Ninad Gujar
  - B. Title: Regulatory Affairs Manager
  - C. Phone Number: 978-564-8503
  - D. FAX Number: 978-750-6677
  - E-Mail: [ngujar@samsungneurologica.com](mailto:ngujar@samsungneurologica.com)
5. **Proposed Device**
  - A. Trade Name: GC85A
  - B. Device Name: GC85A
  - C. Common Name: Digital Diagnostic X-ray System
  - D. Classification Name: System, X-ray, Stationary
  - E. Product Code: KPR
  - F. Regulation: 21 CFR 892.1680
6. **Predicate Device**
  - A. Manufacturer: SAMSUNG ELECTRONICS Co., Ltd.
  - B. Trade Name: XGEO GC80
  - C. Classification Name: System, X-ray, Stationary
  - D. Product Code: KPR
  - E. 510(k) Number: K140334
  - F. 510(k) Decision Date: May 28, 2014
7. **Device Description**

The GC85A digital X-ray imaging system consists of High voltage generator (HVG), Ceiling

510(k) Premarket Notification - Traditional

Suspension, X-ray tube, Collimator, Detector, AEC, DAP, CIB(Control Interface Box), Wall Stand, Patient Table, Collimator, Detector, Remote controller, Grid, Foot switch, Barcode scanner and Auto-stitching stand.

This system is used to capture images by transmitting X-ray to a patient’s body.

The X-ray passing through a patient’s body is sent to the detector and then converted into electrical signals. These signals go through the process of amplification and digital data conversion in the signal process device before being sent to the S-Station (Operation Software) and saved in DICOM file, a standard for medical imaging. The captured images are sent to the Picture Archiving & Communication System (PACS) server, and can be used for reading images.

**8. Intended Use**

The GC85A Digital X-ray Imaging System is intended for use in generating radiographic images of human anatomy by a qualified/trained doctor or technician. This device is not intended for mammographic applications.

**9. Summary of Technological characteristic of the proposed device compared with the predicate device**

The proposed GC85A adds new detectors and modify wall stand to the predicate device, and it does not have significant change in materials, energy source or technological characteristics compared to the predicate device, XGEO GC80 (K140334). Comparisons of the following technological characteristics were assessed and the results demonstrate the substantial equivalence to the predicates.

Specification	Predicate Device	Proposed Device	Discussion
Device Name	XGEO GC80	GC85A	
Manufacturer	SAMSUNG ELECTRONICS	SAMSUNG ELECTRONICS	
510(k) Number	K140334	N/A	
Appearances			Same
Intended Use	The XGEO GC80 digital X-ray imaging system is intended for use in generating radiographic images of human anatomy by a qualified/trained doctor or technician. This device is not intended for mammographic applications.	The GC85A digital X-ray imaging system is intended for use in generating radiographic images of human anatomy by a qualified/trained doctor or technician. This device is not intended for mammographic applications.	Same

Manufacturer Contents	XGEO GC80 (K140334)	GC85A	Discussion
-----------------------	---------------------	-------	------------



510(k) Premarket Notification - Traditional

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion
(1)High Voltage Generator				
Type		High Frequency	High Frequency	Same
Max. Power		80kW	82kW	Difference(1)
Output RANGE	Tube Voltage	40-150kV	40-150kV	Same
	Tube Current	10-1000mA	10-1000mA	Same
	Exposure Time	1msec-6.3sec	1msec-10sec	Difference(2)
AEC (Automatic Exposure Control)		Yes	Yes	Same
APR (Anatomically Programmed Radiography)		Yes	Yes	Same

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion
(2)Ceiling Suspension				
Moving Range (mm)	Longitudinal	1680~4180 (Varies with room size)	1680~4180 (Varies with room size)	Same
	Lateral	1030~3030 (Varies with room size)	1030~3030 (Varies with room size)	Same
	Vertical	1840	1800	Difference(3)
Vertical Tube Moving method		Motorized	Motorized	Same
Tube Assembly Rotation		-157 ~ +183	-157 ~ +183	Same
Brake locking Method		Electromagnetic	Electromagnetic	Same
Automatic Centering		O	O	Same
Moving Rail Type		Al Extrusion	Al Extrusion	Same
Image Preview		O	O	Same
Display Type		Color LCD	Color LCD	Same
Control Switch Type		Button + Touch Screen	Button + Touch Screen	Same
Vertical Sync.	With Table	O	O	Same
	With Stand	O	O	Same

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion
(3) Wall Stand				
Vertical Movement	Mechanism	Motorized/Manual	Motorized	Difference(4)
	Range(mm)	400~1800	280~1850	Difference(5)
Detector/tube servo coupling		Yes	Yes	Same
Detector	Tilting Mechanism	Motorized	Motorized	Same



510(k) Premarket Notification - Traditional

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion
	Range	-20~+90	-20~+90	Same
AEC		Conventional	Conventional	Same
Grid	Lines/cm	84.6	85/92	Difference(6)
	Grid mechanism	Stationary	Stationary	Same
	Removability	Removable	Removable	Same
Detector Support Mounting		Floor	Floor	Same
Patient Support Device		Patient handgrips, lateral support bar	Patient handgrips, lateral support bar	Same

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion	
(4)Patient Table					
Table Top	Size(mm)		2410 X 812	2410 X 812	Same
	Range (mm)	Lateral	±140	±140	Same
		Longitudinal	±480	±480	Same
Table height	Mechanism		DC Motor, Ball screw	DC Motor, Ball screw	Same
	Range(mm)		545 ~ 900	545 ~ 900	Same
Horizontal range of detector(mm)		590	688	Difference(7)	
AEC		Conventional	Conventional	Same	
Grid	Lines/cm		84.6	85/92	Difference(8)
	Grid mechanism		Stationary	Stationary	Same
	Removability		Removable	Removable	Same
Vertical Sync.		O	O	Same	
Control Switch Type		Foot switch	Foot switch	Same	
Maximum Patient Weight(kg)		350 (Static, Center load)	350 (Static, Center load)	Same	

Manufacturer Contents		XGEO GC80 (K140334)	GC85A	Discussion
(5)Collimator				
Overall Size(mm)		H212 X W300 X D179	H212 X W300 X D179	Same
Beam Limiting Blade Moving Method		Motorized /Manual	Motorized /Manual	Same
Manual Operation Method		Volume	Volume	Same
Collimator Rotation		±45	±45	Same
Beam Light Source		LED	LED	Same
Light Field Indicator Timer		O	O	Same
Side Lamp		O	O	Same



510(k) Premarket Notification - Traditional

Manufacturer Contents	XGEO GC80 (K140334)	GC85A	Discussion
	Laser Module	Laser Module	Same
Field Size / SID Display	Color LCD	Color LCD	Same

Manufacturer Contents	XGEO GC80 (K140334)		GC85A			Discussion
(6) Detector						
*NOTE: S4335-W, S4343-W were cleared with K140334.						
Name	S4335-W	S4343-W	S4335-W	S4343-W	S3025-W	
Detector Type	Csl	Csl	Csl	Csl	Csl	Same
	Indirect	Indirect	Indirect	Indirect	Indirect	Same
Detector Area	14"X17" (345mmX42 5mm)	17"X17" (425mmX42 5mm)	14"X17" (345mmX42 5mm)	17"X17" (425mmX42 5mm)	10"X12" (245mmX29 5mm)	Difference(9)
Number of pixels	2466X3040	3036X3040	2466X3040	3036X3040	1750X2108	Difference (10)
Pixel Pitch(um)	140	140	140	140	140	
High Contrast Limiting Resolution (LP/mm)	3.57	3.57	3.57	3.57	3.57	
Communication	Wired / Wireless	Same				

No	Differences	Explanation
(1)	HVG Max Power	Proposed medical device's HVG has higher max power than the predicate device's max power, and the higher max power does not contribute any adverse impacts to the device's safety and performance.
(2)	HVG Exposure Time Range	Proposed medical device's HVG maximum exposure time longer than the predicate device's one, and the longer exposure time does not contribute any adverse impacts to the device's safety and performance.
(3)	Vertical moving range of Ceiling Suspension	Proposed medical device's Vertical moving range of Ceiling Suspension has shorter than the predicate device's moving range, but the shorter moving range does not contribute any adverse impacts to the device's safety and performance.
(4)	Vertical Movement Mechanism of Wall Stand	Proposed medical device's Vertical Movement Mechanism of Wall Stand has a <a href="#">motorized method</a> , while the predicate device's Vertical Movement Mechanism of Wall Stand has <a href="#">two methods as manual &amp; motorized</a> . The movement mechanism does not contribute any adverse impacts to the device's safety and performance.
(5)	Vertical Moving range of	Proposed medical device's Vertical Moving range of Wall Stand has



510(k) Premarket Notification - Traditional

	Wall Stand	longer than the predicate device's one, and the longer moving range does not contribute any adverse impacts to the device's safety and performance.
(6)	Grid line ( Wall stand )	Proposed medical device's Line Pair of Grid installed in Wall Stand has higher than the predicate device's one, and the higher line pair of grid does not contribute any adverse impacts to the device's safety and performance.
(7)	Horizontal range of detector (Patient Table)	Proposed medical device's Horizontal Moving range of Detector installed Patient Table has longer than the predicate device's one, and the longer moving range does not contribute any adverse impacts to the device's safety and performance.
(8)	Grid line (Patient Table)	Proposed medical device's Line Pair of Grid installed in Patient Table has higher than the predicate device's one, and the higher line pair of grid does not contribute any adverse impacts to the device's safety and performance.
(9)	Detector Area	Proposed medical device's S3025-W detector has smaller area than the predicate device's detectors while technical specification is identical among them such as type & pixel pitch, and the smaller area does not contribute any adverse impacts to the device's safety and performance.
(10)	Number of pixels Resolution and pixel pitch of detector	Proposed medical device's S3025-W detector has smaller detector area. Therefore, the proposed medical device's numbers of pixels is smaller than the predicate device's detectors while pixel pitch is identical among them, and the smaller number of pixels does not contribute any adverse impacts to the device's safety and performance.

In non-clinical data, the proposed detectors show curves and measurements of MTF and DQE that do not differ from the predicate device. In clinical data, the proposed GC85A has been shown a substantially equivalent to the predicate device.

**10. Safety, EMC and Performance Data**

Electrical, mechanical, environmental safety and performance testing according to standard ES 60601-1, IEC 60601-1-2, IEC 60601-1-3, IEC 60601-2-28, IEC 60601-2-54, ISO14971, 21CFR1020.30 and 21CFR1020.31 were performed, and EMC testing was conducted in accordance with standard IEC 60601-1-2. Wireless function was tested and verified followed by guidance, Radio frequency Wireless Technology in Medical Devices. All test results were satisfying the standards.

**11. Non-clinical data**

Non-clinical testing data was provided in conformance to the FDA "Guidance for the Submission of 510(k)'s for Solid-State X-ray Imaging Devices", which includes MTF and DQE measurements as tested by IEC 62220-1. The proposed device shows no difference in non-clinical testing data such as MTF and DQE measurements from the predicate device.



**12. Clinical data**

In clinical data, clinical images were obtained in accordance with FDA guidance for the submission of 510(k)'s for Solid State X-ray Imaging Devices. They were evaluated by a professional radiologist and found to be equivalent to the predicate device.

**13. Conclusions**

Clinical images were provided; these images were not necessary to establish substantial equivalence based on the modifications to the device but they provide further evidence in addition to the laboratory performance data to show that the complete system works as intended. These images were evaluated by a radiologist with equivalent U.S. board certification and found to be equivalent to the predicate device.

- 14.** Samsung Electronics Co., Ltd. will update and include in this summary any other information deemed seasonably necessary by the FDA