

Samsung Medison Co., Ltd. % Jee Young Ju Regulatory Affairs Specialist 3366, Hanseo-ro, Nam-myeon Hongcheon-gun, Gangwon-do 25108 REPUBLIC OF KOREA

Re: K220269

Trade/Device Name: HM70 EVO Diagnostic Ultrasound System

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed doppler imaging system

Regulatory Class: Class II Product Code: IYN, IYO, ITX

Dated: March 30, 2022 Received: March 31, 2022

## Dear Jee Young Ju:

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.

May 6, 2022

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 <a href="https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems">https://www.fda.gov/medical-device-problems</a>.

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

Sincerely,

Michael D. O'Hara, Ph.D.

**Deputy Director** 

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

# DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

## Indications for Use

Form Approved: OMB No. 0910-0120 Expiration Date: 06/30/2023

Expiration Date: 06/30/2023 See PRA Statement below.

510(k) Number (if known) K220269
Device Name HM70 EVO Diagnostic Ultrasound System
Indications for Use (Describe) The diagnostic ultrasound system and probes are designed to obtain ultrasound images and analyze body fluids.
The clinical applications include: Fetal/Obstetrics, Abdominal, Gynecology, Pediatric, Small Organ, Neonatal Cephalic, Adult Cephalic, Trans-rectal, Trans-vaginal, Trans-esophageal (Cardiac), Muscular-Skeletal (Conventional, Superficial), Urology, Cardiac Adult, Cardiac Pediatric, Peripheral vessel and Lung.
It is intended for use by, or by the order of, and under the supervision of, an appropriately trained healthcare professional who is qualified for direct use of medical devices. It can be used in hospitals (includes emergency room), private practices, clinics and similar care environment for clinical diagnosis of patients.
Modes of Operation: 2D mode, Color Doppler mode, Power Doppler (PD) mode, M mode, Pulsed Wave (PW) Doppler mode, Continuous Wave (CW) Doppler mode, Tissue Doppler Imaging (TDI) mode, Tissue Doppler Wave (TDW) mode, ElastoScan Mode, Combined modes, Multi-Image mode(Dual, Quad), 3D/4D mode.
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 PRAStaff@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."

# **K220269 - SAMSUNG MEDISON Co., Ltd.**

510(k) Premarket Notification - Traditional

#### 510(k) Summary

In accordance with 21 CFR 807.92 the following summary of information is provided:

1. Date Prepared – January 28, 2022

2. Manufacturer

SAMSUNG MEDISON CO., LTD. 3366, Hanseo-ro, Nam-myeon, Hongcheon-gun, Gangwon-do 25108, REPUBLIC OF KOREA

3. Primary Contact Person

Jee Young Ju

Regulatory Affairs Specialist

Phone: +82.2.2194.0861 Fax: +82.2.2194.0278

Email: jee.ju@samsungmedison.com

4. Secondary Contact Person

Ninad Gujar Vice President

Phone: +1.978.564.8632 Fax: +1.978.564.8677

Email: ngujar@neurologica.com

5. Proposed Device

- Proprietary Name: HM70 EVO Diagnostic Ultrasound System - Common Name: System, Imaging, Pulsed Doppler, Ultrasonic System, Imaging, Pulsed Echo, Ultrasonic

Transducer, Ultrasonic, Diagnostic

- Classification: 21 CFR 892.1550 Ultrasonic pulsed doppler imaging system

21 CFR 892.1560 Ultrasonic pulsed echo imaging system

21 CFR 892.1570 Diagnostic ultrasonic transducer

- Product Code(s): IYN, IYO, ITX

6. Predicate Device

HM70 EVO Diagnostic Ultrasound System (K210713) - Primary Predicate HM70A Diagnostic Ultrasound System (K182894) - Reference V8 Diagnostic Ultrasound System (K211945) - Reference

#### 7. Device Description

The HM70 EVO is a general purpose, mobile, software controlled, diagnostic ultrasound system. Its function is to acquire ultrasound data and to display the data as 2D mode, M mode, Color Doppler imaging, Power Doppler imaging (including Directional Power Doppler mode; S-Flow), PW Spectral Doppler mode, CW Spectral Doppler mode, Harmonic imaging(S-Harmonic), Tissue Doppler imaging, Tissue Doppler Wave, Panoramic Imaging, Freehand 3D, 3D imaging mode (real-time 4D imaging mode), Elastoscan Mode or as a combination of these modes. The HM70 EVO also gives the operator the ability to measure

# SAMSUNG MEDISON Co., Ltd.

510(k) Premarket Notification – Traditional

anatomical structures and offers analysis packages that provide information that is used to make a diagnosis by competent health care professionals. The HM70 EVO has real time acoustic output display with two basic indices, a mechanical index and a thermal index, which are both automatically displayed.

#### 8. Indications for Use

The diagnostic ultrasound system and probes are designed to obtain ultrasound images and analyze body fluids.

The clinical applications include: Fetal/Obstetrics, Abdominal, Gynecology, Pediatric, Small Organ, Neonatal Cephalic, Adult Cephalic, Trans-rectal, Trans-vaginal, Trans-esophageal (Cardiac), Muscular-Skeletal (Conventional, Superficial), Urology, Cardiac Adult, Cardiac Pediatric, Peripheral vessel and Lung.

It is intended for use by, or by the order of, and under the supervision of, an appropriately trained healthcare professional who is qualified for direct use of medical devices. It can be used in hospitals (includes emergency room), private practices, clinics and similar care environment for clinical diagnosis of patients.

Modes of Operation: 2D mode, Color Doppler mode, Power Doppler (PD) mode, M mode, Pulsed Wave (PW) Doppler mode, Continuous Wave (CW) Doppler mode, Tissue Doppler Imaging (TDI) mode, Tissue Doppler Wave (TDW) mode, ElastoScan Mode, Combined modes, Multi-Image mode(Dual, Quad), 3D/4D mode.

## 9. Technology

The HM70 EVO employs the same fundamental scientific technology as its predicate devices.

#### 10. Determination of Substantial Equivalence

Comparison to Predicates: The HM70 EVO is substantially equivalent to the predicate devices with regard to intended use, imaging capabilities, technological characteristics and safety and effectiveness.

- The systems are all intended for diagnostic ultrasound imaging and fluid flow analysis
- The proposed HM70 EVO and predicates HM70 EVO(K210713), HM70A (K182894) and V8(K211945) have the same clinical intended use, imaging modes and modes of operation.
- The proposed HM70 EVO has added the LS6-15 already cleared in the predicate device HM70A(K182894), PA1-5A & MMPT3-7 already cleared in the predicate device V8(K211945) and the new transducers LA2-9S, PA3-9B.
- The proposed HM70 EVO has added the 2D Follicle and Mobile Export already cleared in V8(K211945)
- The system is manufactured with materials which have been evaluated and found to be safe for the intended use of the device.

# SAMSUNG MEDISON Co., Ltd.

510(k) Premarket Notification – Traditional

- The system has acoustic power levels which are below the applicable FDA limits.
- The proposed HM70 EVO and predicate HM70 EVO(K210713), HM70A(K182894) and V8(K211945) have similar capability in terms of performing measurements, capturing digital images, reviewing and reporting studies.
- The proposed HM70 EVO and predicates have been designed in compliance with approved electrical and physical safety standards.

#### 11. Summary of Non-Clinical Test

The device has been evaluated for acoustic output, biocompatibility, software function, cleaning and disinfection effectiveness as well as thermal, electrical, electromagnetic and mechanical safety, and has been found to conform with applicable FDA guidance and medical device safety standards. The HM70 EVO and its applications comply with the following FDA-recognized standards.

Reference No.	Title
IEC 60601-1	AAMI ANSI ES60601-1:2005/(R)2012 and A1:2012, C1:2009/(R)2012 and A2:2010/(R)2012 (Consolidated Text) Medical electrical equipment -
	Part 1: General requirements for basic safety and essential performance (IEC 60601-1:2005, MOD)
IEC 60601-1-2	IEC60601-1-2: 2014(4th Edition), Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - EMC
	IEC 60601-2-18: Edition 3.0 2009-08
IEC 60601-2-18	Medical electrical equipment - Part 2-18: Particular requirements for the
	basic safety and essential performance of endoscopic equipment
IEC 60601-2-37	IEC 60601-2-37 Edition 2.0 2007, Medical electrical equipment – Part 2-
	37: Particular requirements for the basic safety and essential performance
	of ultrasonic medical diagnostic and monitoring equipment
ISO10993-1	ISO 10993-1:2009/(R)2013, Biological evaluation of medical devices –
	Part 1: Evaluation and testing within a risk management process
ISO14971	ISO 14971:2007, Medical devices - Application of risk management to
	medical devices
NEMA UD 2-2004	NEMA UD 2-2004 (R2009)
	Acoustic Output Measurement Standard for Diagnostic Ultrasound
	Equipment Revision 3

#### 12. Summary of Clinical Tests

The subject of this premarket submission, HM70 EVO, is not required clinical studies to support substantial equivalence.

#### 13. Conclusion

Since the predicates device and subject device have a similar intended use and key technological features, the non-clinical data support the safety of the device and demonstrate that the HM70 EVO Ultrasound System should perform as intended in the specified use conditions. Therefore, SAMSUNG MEDISON CO., LTD. considers the HM70 EVO to be as safe, as effective, and provide performance substantially equivalent to the primary predicate device(K210713).

## - END of 510(k) Summary -