



April 13, 2018

Edan Instruments, Inc.
% Mr. Doug Worth
Sr. Dir. US RA/QA
Edan Medical
1200 Crossman Avenue, Suite 200
SUNNYVALE CA 94089

Re: K180408

Trade/Device Name: U60 Diagnostic Ultrasound System
Regulation Number: 21 CFR 892.1550
Regulation Name: Ultrasonic pulsed doppler imaging system
Regulatory Class: II
Product Code: IYN, IYO, ITX
Dated: March 28, 2018
Received: April 2, 2018

Dear Mr. Worth:

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.

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.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<https://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/>) and CDRH Learn (<http://www.fda.gov/Training/CDRHLearn>). 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 (<http://www.fda.gov/DICE>) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,



Robert Ochs, Ph.D.
Director
Division of Radiological Health
Office of In Vitro Diagnostics
and Radiological Health
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

Form Approved: OMB No. 0910-0120
Expiration Date: January 31, 2017
See PRA Statement below.

Indications for Use

510(k) Number (if known)

K180408

Device Name

U60 Diagnostic Ultrasound System

Indications for Use (Describe)

The diagnostic ultrasound system (U60) is applicable for adults, pregnant women, pediatric patients' ultrasound evaluation in hospitals and clinics. It is intended for use in abdominal, obstetrics, gynecology, pediatric, small parts, urology, peripheral vascular, musculoskeletal (conventional and superficial), endovaginal and cardiac clinical applications, by or on the order of a physician or similarly qualified health care professional.

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

"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."

Diagnostic Ultrasound Indications for Use Form

U60 Diagnostic Ultrasound System

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics	P	P	P		P	P	P
	Abdominal	P	P	P		P	P	P
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric	P	P	P		P	P	P
	Small Organ (Specify) *	P	P	P		P	P	P
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal	P	P	P		P	P	P
	Trans-urethral							
	Musculo-skeletal(Conventional)	P	P	P		P	P	P
	Musculo-skeletal (Superficial)	P	P	P		P	P	P
	Intravascular							
Other (Specify) **	P	P	P		P	P	P	
Cardiac	Adult Cardiac	P	P	P	P	P	P	P
	Pediatric Cardiac	P	P	P		P	P	P
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular	P	P	P		P	P	P
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]:PDI: Power Doppler Imaging , DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with C352UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics	P	P	P		P	P	P
	Abdominal	P	P	P		P	P	P
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **	P	P	P		P	P	P	
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with L1042UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *	P	P	P		P	P	P
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)	P	P	P		P	P	P
	Musculo-skeletal (Superficial)	P	P	P		P	P	P
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular	P	P	P		P	P	P
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with L742UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *	P	P	P		P	P	P
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)	P	P	P		P	P	P
	Musculo-skeletal (Superficial)	P	P	P		P	P	P
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular	P	P	P		P	P	P
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with E612UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics	P	P	P		P	P	P
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal	P	P	P		P	P	P
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with C612UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric	P	P	P		P	P	P
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac	P	P	P		P	P	P
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with C6152UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric	P	P	P		P	P	P
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac	P	P	P		P	P	P
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with C422UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal	P	P	P		P	P	P
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac	P	P	P		P	P	P
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with L552UB Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric	P	P	P		P	P	P
	Small Organ (Specify) *	P	P	P		P	P	P
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)	P	P	P		P	P	P
	Musculo-skeletal (Superficial)	P	P	P		P	P	P
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular	P	P	P		P	P	P
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with C5-2b Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

General	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics	P	P	P		P	P	P
	Abdominal	P	P	P		P	P	P
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **		P	P	P		P	P	P
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with P5-1b Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *							
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)							
	Musculo-skeletal (Superficial)							
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac	P	P	P	P	P	P	P
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular							
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

Diagnostic Ultrasound Indications for Use Form

U60 with L15-7b Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application	Mode of Operation						
		B	M	PW	CW	Color	Combined (Specify) ^[1]	Other (Specify) ^{[2][3]}
General	Specific							
Ophthalmic	Ophthalmic							
Fetal Imaging & Other	Fetal / Obstetrics							
	Abdominal							
	Intra-operative (Specify)							
	Intra-operative (Neuro logical)							
	Laparoscopic							
	Pediatric							
	Small Organ (Specify) *	P	P	P		P	P	P
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							
	Musculo-skeletal(Conventional)	P	P	P		P	P	P
	Musculo-skeletal (Superficial)	P	P	P		P	P	P
	Intravascular							
Other (Specify) **								
Cardiac	Adult Cardiac							
	Pediatric Cardiac							
	Intravascular(Cardiac)							
	Trans-esoph.(Cardiac)							
	Intra- cardiac							
Peripheral vascular	Peripheral vascular	P	P	P		P	P	P
	Other (Specify)							

N = new indication; P = previously cleared by FDA; E = added under this appendix PDI= Power Doppler Imaging

Additional comments: Combined mode: B+M, B+PW, B+Color, B+PDI/DPDI, B+Color+PW, B+PDI/DPDI +PW

Note * Small Organ includes Thyroid, Testes, Breast

** Other use includes Urology, Gynecology

[1]: PDI: Power Doppler Imaging ,DPDI: Directional Power Doppler Imaging

[2]: Biopsy Guidance

[3]: Harmonic Imaging, This feature does not use contrast agent.

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Device Evaluation and Safety (OIVD)

Prescription Use (Per 21 CFR 801.109)

510(k) Summary

Prepared in accordance with the requirements of 21 CFR Part 807.92

- 1. Submitter:** Edan Instruments, Inc.
#15 Jinhui Road, Jinsha Community,
Kengzi Sub-District, Pingshan District, Shenzhen, 518122 P.R.China.
Tel.: (0755) 26858736
Fax: +1 (408) 418-4059
- Contact Person:** Crystal Cai
- Date prepared:** February 12, 2018
- 2. Device name and classification:** **Device Name:** U60 Diagnostic Ultrasound System
Model: U60
Classification Name:
892.1550 System, Imaging, Pulsed Doppler, Ultrasonic
Product code: IYN
892.1560 Ultrasonic, Pulsed echo, Imaging
Product code: IYO
892.1570 Transducer, Ultrasonic, Diagnostic
Product code: ITX
Regulatory Class: Class II
- 3. Premarket Notification Class III Certification and Summary** Not applicable, the subject device is Class II.
- 4. Predicate Device(s):** U50 Diagnostic ultrasound system/ K173003/ Edan Instruments, Inc.
- 5. Pre-Submission, IDE** Not applicable, there is no prior submission.

6. Device Description:

The U60 is a portable Diagnostic Ultrasound System, which applies advanced technologies. Various image parameter adjustments, 15.0 inch LCD and diverse probes are configured to provide clear and stable images.

7. Intended Use:

The diagnostic ultrasound system (U60) is applicable for adults, pregnant women, pediatric patients' ultrasound evaluation in hospitals and clinics. It is intended for use in abdominal, obstetrics, gynecology, pediatric, small parts, urology, peripheral vascular, musculoskeletal (conventional and superficial), endovaginal and cardiac clinical applications, by or on the order of a physician or similarly qualified health care professional.

9. Predicate Device Comparison

Comparison to the predicate devices, the subject device has the same intended use, similar product design, same performance effectiveness, performance safety as the predicate device as summarized in the following tables:

Item	U60 R1.0 Diagnostic Ultrasound System (Edan Instruments)	U50 R2.2 Diagnostic Ultrasound System (Edan Instruments)	Comparison Result
510(k) Number	Current Submission	K173003	
Manufacturer	EDAN Instruments	EDAN Instruments	Same
Intended Use	Diagnostic ultrasound imaging or fluid flow analysis of the human body	Diagnostic ultrasound imaging or fluid flow analysis of the human body	Same
Indications for Use	The diagnostic ultrasound system (U60) is applicable for adults, pregnant women, pediatric patients' ultrasound evaluation in hospitals and clinics. It is intended for use in abdominal, obstetrics, gynecology, pediatric, small parts, urology, peripheral vascular, musculoskeletal (conventional and superficial), endovaginal and cardiac clinical applications, by or on the order of a physician or similarly qualified health care professional.	The diagnostic ultrasound system (U50) is applicable for adults, pregnant women, pediatric patients' ultrasound evaluation in hospitals and clinics. It is intended for use in abdominal, obstetrics, gynecology, pediatric, small parts, urology, peripheral vascular, musculoskeletal (conventional and superficial), endovaginal and cardiac clinical applications, by or on the order of a physician or similarly qualified health care professional.	Same
Installation and Use	a. Portable Equipment b. Mobile Equipment (when the system is installed on the mobile trolley)	a. Portable Equipment b. Mobile Equipment (when the system is installed on the mobile trolley)	Same

Safety Standards	IEC 60601-1 IEC 60601-1-2 IEC 60601-2-37 ISO 10993-1, -5, -10 AIUM, NEMA UD 2, UD3	IEC 60601-1 IEC 60601-1-2 IEC 60601-2-37 ISO 10993-1, -5, -10 AIUM, NEMA UD 2, UD3	Same
Patient Contact Materials	Complies with ISO 10993	Complies with ISO 10993	Same
Software life cycle processes	Complies with the standard: IEC 62304	Complies with the standard: IEC 62304	Same
Mode of Operations	Continuous operation	Continuous operation	Same
General Imaging mode	B-Mode, M-Mode, Color, PDI/DPDI, PW, CW	B-Mode, M-Mode, Color, PDI/DPDI, PW, CW	Same
Measurements	B-Mode: Distance, Area, Volume, Ratio, Histogram and Angle M-Mode: Distance, Time, Slope, and Heart Rate D-Mode: Time, Heart Rate, Velocity, Acceleration, RI, PI and Auto (auto trace)	B-Mode: Distance, Area, Volume, Ratio, Histogram and Angle M-Mode: Distance, Time, Slope, and Heart Rate D-Mode: Time, Heart Rate, Velocity, Acceleration, RI, PI and Auto (auto trace)	Same
Scanning method	Electronic convex Electronic linear with slant scanning	Electronic convex Electronic linear with slant scanning	Same
Cine loop	1227 frames	1227 frames	Same
Focus Number	Max=4	Max=4	Same
Software Packages	Abdomen, obstetric, small parts, gynecology, cardiology, urology, vascular and Pediatrics.	Abdomen, obstetric, small parts, gynecology, cardiology, urology, vascular and Pediatrics.	Same
Principle of Operation	Applying high voltage burst to the Piezoelectric material in the transducer and detect reflected echo to construct diagnostic image	Applying high voltage burst to the Piezoelectric material in the transducer and detect reflected echo to construct diagnostic image	Same
Acoustic Output	Track 3: MI, TIS, TIC, TIB(TI Range 0-6.0) Derated I_{SPTA} : 720 W/cm ² maximum, Mechanic Index ≤ 1.9 maximum or Derated I_{SPPA} 190 W/cm ² max	Track 3: MI, TIS, TIC, TIB(TI Range 0-6.0) Derated I_{SPTA} : 720 W/cm ² maximum, Mechanic Index ≤ 1.9 maximum or Derated I_{SPPA} 190 W/cm ² max	Same
Transducer Types	Convex Array Linear Array Micro Convex Array Phased Array	Convex Array Linear Array Micro Convex Array Phased Array	Same
Transducer Frequency	2.0-15.0 MHz	2.0-15.0 MHz	Same
Primary Display	Primary Screen: 15inch(1024*768)	Primary Screen: 12.1inch(1024*768)	Difference
Transducer	Multi-Transducer Port (Two)	Multi-Transducer Port (Two)	Same

Ports			
Dimensions/ Weight	220 mm (W) × 370 mm (L) × 350 mm (H) net weight 8.1kg	330 mm (W) × 320 mm (L) × 220 mm (H) net weight 7.8kg	Difference
Printer	B/W video thermal printer Color video thermal printer Graph/text laser jet printer	B/W video thermal printer Color video thermal printer Graph/text laser jet printer	Same
Storage Media	DVD, USB stick	DVD, USB stick	Same
Temperature	Operating: 0°C ~40°C Transport/ Storage: -20~55°C	Operating: 5°C ~40°C Transport/ Storage: -20~55°C	Difference
Relative humidity	Operating: 15% ~ 95%RH (no condensation) Transport/ Storage: 15% ~ 95%RH (no condensation)	Operating: 25% ~ 80%RH (no condensation) Transport/ Storage: 25% ~ 93%RH (no condensation)	Difference
Atmospheric pressure	Operating: 860hPa ~1060hPa Transport/ Storage: 700hPa ~1060hPa	Operating: 860hPa ~1060hPa Transport/ Storage: 700hPa ~1060hPa	Same
Power Requirements	AC: 100-240V 50/60Hz	AC: 100-240V 50/60Hz	Same
Operation System	Linux	Linux	Same
Safety Classifications			
Type of protection against electric shock	Class I	Class I	Same
The degree of protection against electric shock	Type BF.	Type BF.	Same
The degree of protection against harmful ingress of liquid	The main unit : IPX0, probes : IPX7 Footswitch: IP68	The main unit : IPX0, probes : IPX7 Footswitch: IP68	Same
The degree of safety of application in the presence of a flammable gas	Equipment not suitable for use in the presence of a flammable gas	Equipment not suitable for use in the presence of a flammable gas	Same
The degree of RF	Group 1, Class A	Group 1, Class A	Same
Disinfection			
Disinfection	Probe: 2.4%Glutaraldehyde, 0.55%Ortho-Phthalaldehyde. Needle guide: 75% medical alcohol,	Probe: 2.4%Glutaraldehyde, 0.55%Ortho-Phthalaldehyde. Needle guide: 75% medical alcohol,	Same

	2.4%Glutaraldehyde.	2.4%Glutaraldehyde.	
Performance			
Displayed depth	20-320mm (Probe Dependent)	20-320mm (Probe Dependent)	Same
Gray Scales	256	256	Same
Dynamic range	150dB	150dB	Same
TGC	8 segments	8 segments	Same
Zoom	Up to 400%	Up to 400%	Same
Image Adjustments			
B Mode Parameters	Image type	Image type	Same
	Gain	Gain	
	Depth	Depth	
	TGC	TGC	
	Freq(Frequency)	Freq(Frequency)	
	Gray Map	Gray Map	
	Dynamic Range	Dynamic Range	
	Rejection	Rejection	
	Focus Position	Focus Position	
	Focus Number	Focus Number	
	eSRI(Spackle Imaging) Rejection	eSRI(Spackle Imaging) Rejection	
	Pseudo color	Pseudo color	
	Spatial Compound	Spatial Compound	
	GAO(Gain Auto Optimization)	GAO(Gain Auto Optimization)	
	Scan Angle	Scan Angle	
	Scan Mode	Scan Mode	
	Frame Persist	Frame Persist	
	H Reverse(horizontal)	H Reverse(horizontal)	
V Reverse(vertical)	V Reverse(vertical)		
90°Rotate	90°Rotate		
B/W Invert	B/W Invert		
M Mode Parameters	Freq	Freq	Same
	Sweep Speed	Sweep Speed	
	Display Layout	Display Layout	
	Gray Map	Gray Map	
	Focus Position	Focus Position	
	Dynamic Range	Dynamic Range	
	Pseudo Color	Pseudo Color	
Line Average	Line Average		
Color Mode & Power Doppler Mode & Directional Power Mode Parameters	Flow type	Flow type	Same
	Gain	Gain	
	Freq	Freq	
	Wallfilter	Wallfilter	
	PRF(Pulsed Repetition Frequency)	PRF(Pulsed Repetition Frequency)	
	Base Line	Base Line	
	Invert	Invert	
	Dual Live	Dual Live	
Angle Steer	Angle Steer		
Color Map	Color Map		

	Packet size	Packet size	
	Persist	Persist	
	Threshold	Threshold	
	Smooth Filter	Smooth Filter	
	ROI box position and size adjustment	ROI box position and size adjustment	
PW Mode Parameters	Flow type	Flow type	Same
	Gain	Gain	
	PRF	PRF	
	Invert	Invert	
	Angle steer	Angle steer	
	Correction Angle	Correction Angle	
	Quick Angle	Quick Angle	
	Base Line	Base Line	
	Sample Volume	Sample Volume	
	Wallfilter	Wallfilter	
	Freq	Freq	
	Duplex and Triplex	Duplex and Triplex	
	Pseudo Color	Pseudo Color	
	Dyn Rng	Dyn Rng	
	Volume	Volume	
Sweep Speed	Sweep Speed		
HPRF	HPRF		
CW Mode Parameters	Flow type	Flow type	Same
	Gain	Gain	
	PRF	PRF	
	Invert	Invert	
	Angle steer	Angle steer	
	Correction Angle	Correction Angle	
	Quick Angle	Quick Angle	
	Base Line	Base Line	
	Wallfilter	Wallfilter	
	Pseudo Color	Pseudo Color	
	Dynamic Range	Dynamic Range	
	Volume	Volume	
	Sweep Speed	Sweep Speed	
B-Mode Measurement Accuracy			
Range of Depth/Distance	Maximum 324 mm	Maximum 324 mm	Same
Accuracy of Depth/Distance	$\leq \pm 5\%$	$\leq \pm 5\%$	Same
Range of Area	Maximum 1126 cm ²	Maximum 1126 cm ²	Same
Accuracy of Area	$\leq \pm 10\%$	$\leq \pm 10\%$	Same
Range of Angle	0-180°	0-180°	Same
Accuracy of Angle	$\leq \pm 3\%$	$\leq \pm 3\%$	Same
Range of	Maximum 1.0	Maximum 1.0	Same

Ratio			
Accuracy of Ratio	≤±10%	≤±10%	Same
Range of Volume	Maximum 999 cm ³	Maximum 999 cm ³	Same
Accuracy of Volume	≤±15%	≤±15%	Same
M-mode Measurement Accuracy			
Range of Depth	Maximum 324mm	Maximum 324mm	Same
Accuracy of Depth	≤±5%	≤±5%	Same
Range of Time	Maximum 13s	Maximum 13s	Same
Accuracy of Time	≤±5%	≤±5%	Same
Range of Heart rate	Maximum 999bpm	Maximum 999bpm	Same
Accuracy of Heart rate	≤±5%	≤±5%	Same
Range of Slope	Maximum 999mm/s	Maximum 999mm/s	Same
Accuracy of Slope	≤±10%	≤±10%	Same
PW mode velocity Measurement Accuracy			
Range	0.5-2.5m/s	0.5-2.5m/s	Same
Accuracy	≤±10%	≤±10%	Same
CW mode velocity Measurement Accuracy			
Range	0.5-2.5m/s	0.5-2.5m/s	Same
Accuracy	≤±10%	≤±10%	Same

The subject device has same intended use, similar product design, and same performance effectiveness, as the predicate device. There are no significant differences between the primary predicate. There are no new questions of safety and/or effectiveness. There are no changes to the intended use, indications for use, nor fundamental scientific technology.

10. Effectiveness and Safety Considerations:

Clinical test:

Clinical testing is not required.

Non-clinical test:

The U60 Diagnostic Ultrasound System complies with:

- (1) ANSI AAMI 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).

- (2) IEC 60601-1-2:2014: Medical electrical equipment - part 1-2: general requirements for basic safety and essential performance - collateral standard: electromagnetic compatibility - requirements and tests.
- (3) 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.
- (4) NEMA UD 3, Edition 2004 Standard for real-time display of thermal and mechanical acoustic output Indies on diagnostic ultrasound equipment.
- (5) Acoustic output testing as per the guideline “Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers” dated September 9, 2008.
- (6) NEMA, UD2, Edition 2004 for acoustic output measurement methodology.

The following biocompatibility standards are conducted on the subject device:

- (1) ISO 10993-1:2009, ISO 10993-5:2009 and ISO 10993-10:2010

The tests were selected to show substantial equivalence between the subject device and the predicate.

11. Substantially Equivalent Determination

Verification and validation testing has been conducted on the U60 Ultrasound Imaging System. This premarket notification submission demonstrates that U60 Ultrasound Imaging System is substantially equivalent to the predicate devices.