KOZ1118

DEC 2 2 2008

510(k) Summary Prepared August 27, 2008 Revised December 18, 2008

Sponsor:	Siemens Medical Solutions USA, Inc.,
	Ultrasound Division
	1230 Shorebird Way
	P.O. Box 7393
	Mountain View, California 94039-7393

Contact Person:	Sheila W. Pick	ering Ph.D.
	Telephone:	(650) 943 7187
	Fax:	(650) 943 7053

Submission Date:	April 18, 2008

Device Name: Acuson S2000 ABVS Ultrasound System

Common Name: Diagnostic Ultrasound System with Accessories

Classification:

Regulatory Class: II Review Category: Tier II Classification Panel: Radiology

Ultrasonic Pulsed Doppler Imaging System	FR # 892.1550	Product Code 90-IYN
Ultrasonic Pulsed Echo Imaging System	FR # 892.1560	Product Code 90-IYO
Diagnostic Ultrasound Transducer	FR # 892.1570	Product Code 90-ITX

A. Legally Marketed Predicate Devices

The Acuson S2000 ABVS Ultrasound system is substantially equivalent to the Acuson S2000 ultrasound system (K072786) and the U-Systems ultrasound system.

B. Device Description:

The Acuson S2000 ABVS Ultrasound System has been designed to meet the following product safety standards:

- UL 60601-1, Safety Requirements for Medical Equipment
- IEC 60601-2-37 Diagnostic Ultrasound Safety Standards
- CSA C22.2 No. 601-1, Safety Requirements for Medical Equipment
- AIUM/NEMA UD-3, 1998 Standard for Real Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment
- AIUM/NEMA UD-2, 1998 Acoustic Output Measurement Standard for Diagnostic Ultrasound
- 93/42/EEC Medical Devices Directive
- Safety and EMC Requirements for Medical Equipment
 - EN/IEC 60601-1
 - EN/IEC 60601-1-1
 - EN/IEC 60601-1-2
- IEC 1157 Declaration of Acoustic Power
- ISO 10993-1 Biocompatibility

C. Intended Use

The Modified S2000, the S2000 ABVS Ultrasound System, is intended for the following applications: General Radiology, Abdominal, Fetal, Small Parts, Transcranial, OB/GYN, Cardiac, Pelvic, Neonatal/Adult Cephalic, Pediatric, Urology, Vascular, Musculoskeletal, Superficial Musculoskeletal, and Peripheral Vascular applications. The system supports the transducers listed in the Ultrasound Indications for Use tables, including the 14L5BV for B-mode imaging of a patient's breast using an optional automatic scanning function. The device is not intended to be used as a replacement for screening mammography.

The system also provides for the measurement of anatomical structures and analysis as provided in the original S2000 Ultrasound System, which provides information that is used by medical health care professionals for clinical diagnosis purposes.

D. Substantial Equivalence

The submission device is substantially equivalent to the predicate devices with regard to both intended use and technological characteristics.

E. Performance Data

The S2000 modifications are verified and validated according to the company's design control process.



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

Sheila W. Pickering, Ph.D. Senior Director of Regulatory Affairs Siemens Medical Solutions USA, Inc. Ultrasound Division 1230 Shorebird Way MOUNTAIN VIEW WAY CA 94039-7393

DEC 2 2 2008

Re: K081148

Trade/Device Name: Acuson S2000 ABVS Ultrasound System Regulation Number: 21 CFR 892.1550 Regulation Name: Ultrasonic pulsed doppler imaging system Regulatory Class: II Product Code: IYN, IYO, and ITX Dated: November 21, 2008 Received: November 26, 2008

Dear Dr. Pickering:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and we 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). 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.

This determination of substantial equivalence applies to the following transducers intended for use with the Acuson S2000 ABVS Ultrasound System, as described in your premarket notification:

Transducer Model Number

<u>CW2 Probe</u> <u>CW5 Probe</u> <u>EC9-4 Curved Array</u> <u>9L4 Linear Array</u> <u>14L5 Multi-D Array</u> <u>4P1 Phased Array</u> <u>6C2 Curved Array</u> <u>4C1 Curved Array</u> <u>4V1 Phased Array</u>

<u>10V4 Phased Array</u> <u>14L5 SP Linear Array</u> <u>7CF2 Curved Array Mechanical 3D</u> <u>9EVF4 Curved Array</u> <u>V5Ms Multiplane TEE</u> <u>18L6HD Linear Array</u> <u>8V3 Phased Array</u>

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such 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); 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.

This letter will allow you to begin marketing your device as described in your premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus permits your device to proceed to market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (240) 276-3150 or at its Internet address http://www.fda.gov/cdrh/industry/support/index.html

If you have any questions regarding the content of this letter, please contact Lauren Hefner at (240) 276-3666.

Sincerely yours

Joyce M. Whang, Ph.D. Acting Director, Division of Reproductive, Abdominal, and Radiological Devices Office of Device Evaluation Center for Devices and Radiological Health

Enclosure(s)

Indications for Use

510(k) Number (if known): K081148

Device Name:

Acuson S2000 ABVS Ultrasound System

Indications For Use:

The Modified S2000, the S2000 ABVS Ultrasound System, is intended for the following applications: General Radiology, Abdominal, Fetal, Small Parts, Transcranial, OB/GYN, Cardiac, Pelvic, Neonatal/Adult Cephalic, Pediatric, Urology, Vascular, Musculoskeletal, Superficial Musculoskeletal, and Peripheral Vascular applications. The system supports the transducers listed in the Ultrasound Indications for Use tables, including the 14L5BV for B-mode imaging of a patient's breast using an optional automatic scanning function. The device is not intended to be used as a replacement for screening mammography.

The system also provides for the measurement of anatomical structures and analysis as provided in the original S2000 Ultrasound System, which provides information that is used by medical health care professionals for clinical diagnosis purposes.

Prescription Use X AND/OR (Part 21 CFR 801 Subpart D)

<u>بن</u>

Over-The-Counter Use _____(21 CFR 801 Subpart C)

(PLE	ASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER
PAGE 🎼 NI	EEDED)
	s Keyn
(Division Sign-Off)	Concurrence of CDRH, Office of Device Evaluation (ODE)
Division of Reproductiv	e, Abdominal and
Radiological Devices	1/11/0
510(k) Number	KO8 1148

Prescription Use (Per 21 CFR 801.10 Diagnostic Ultrasound Indications for Use Form

510 (k) Number (if known):

Device Name:

CW2 Probe for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation									
Clinical Application	А	в	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)
Ophthalmic										
Fetal			····		P					
Abdominal				[P					
Intraoperative (Note 9)					Р					
Intraoperative Neurological										
Pediatric					P					
Small Organ (Note 1)				:	P					
Neonatal Cephalic					Р					
Adult Cephalic					Р					
Cardiac					P					
Trans- esophageal		l.								
Transrectal										
Transvaginal										
Transurethral										
Intravascular					l					
Peripheral vessel					P_					
Laparoscopic									ļ	
Musculo-skeletal Conventional					P					
Musculo-skeletal Superficial					P					
Other (specify)		1					1			

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc. Note 9 For example: vascular, abdominal

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF
NEEDED)
Concurrence of CDRH, Office of Device Evaluation (ODE)
Prescription Use (Per 21 CFR 801.109)
(Division Sign-Off)
Division of Reproductive, Abdominal and
Radiological Devices $1/\sqrt{2}/14$
510(k) Number 0 0// (8

510 (k) Number (if known):

K 58/148

Device Name:

CW5 Probe for use with ACUSON S2000 ABVS Ultrasound System Ultrasound imaging or fluid flow analysis of the human body as follows:

Intended Use:

Mode of Operation Color Amplitud Combin Color Other Clinical PWDICWD Velocity ed А в Μ е (Specify) Doppler Application Imaging (Specify) Doppler **Ophthalmic** Ρ Fetal P Abdominal Intraoperative Ρ (Note 9) Intraoperative Neurological Ρ Pediatric Small Organ Ρ (Note 1) Neonatal Ρ Cephalic P Adult Cephalic Ρ Cardiac Transesophageal Transrectal Transvaginal Transurethral Intravascular Ρ Peripheral vessel Laparoscopic Musculo-skeletal Ρ Conventional Musculo-skeletal Ρ Superficial Other (specify)

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

510(k) Number _

K081148

Note 1 For example: breast, testes, thyroid, penis, prostate, etc. Note 9 For example: vascular, abdominal

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)
Concurrence of CDRH, Office of Device Evaluation (ODE)
Prescription Use (Per 21 CFR 801.109)
(Division Sign-Off)
Division of Reproductive, Abdominal and
Radiological Devices

510 (k) Number (if known):

K081148

Device Name:

EC9-4 Curved Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation									
Clinical Application	А	B	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Ρ	Ρ	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11	
Abdominal		P	Ρ	Р		Р	Р		BMDC	Note 2,3,4,5,6,,7,8,10, 11	
Intraoperative Abdominal											
Intraoperative Neurological											
Pediatric		ļ								Note	
Small Organ (Note 1)	-	Р	Р	P		P	Р		BMDC	2,3,4,5,7,8,10, 11,14	
Neonatal Cephalic		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10	
Adult Cephalic											
Cardiac									<u> </u>		
Trans- esophageal											
Transrectal		Р	P	P.		Р	P		BMDC	Note 2,3,4,5,7,8,10, 11,14	
Transvaginal		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11	
Transurethral											
Intravascular						1					
Peripheral vessel			·		ļ	ļ	<u></u>		<u> </u>		
Laparoscopic					ļ	<u> </u> .	<u> </u>	L			
Musculo-skeletal Conventional											
Musculo-skeletal Superficial											
Other (specify)									<u> </u>		

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc.

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

(Division Sign-Off)	-
Division of Reproductive Abdomination	
510(k) NumberK081148	

- Note 4 Tissue Equalization Technology
- Note 5 3-Scape real-time 3D imaging
- Note 6 Cadence contrast agent imaging
- Note 7 B&W SieScape panoramic imaging
- Note 8 Power SieScape panoramic imaging
- Note 10 Clarify VE vascular enhancement technology
- Note 11 Advanced Sieclear multi-view spatial compounding
- Note 14 eSie™ Touch elasticity imaging/FTI

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

10 .

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number _____K 08/14/8

510 (k) Number (if known):

<08/148

Device Name:

9L4 Linear Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation									
Clinical Application	А	в	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Ρ	Ρ	Ρ		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11	
Abdominal Intraoperative Abdominal											
Intraoperative Neurological					,						
Pediatric		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11	
Small Organ (Note 1)		Р	P	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14	
Neonatal Cephalic		Р	Р	Р		Р	Р	· · ·	BMDC	Note 2,3,4,5,7,8,10, 11	
Adult Cephalic Cardiac			<u> </u>							·····	
Trans- esophageal											
Transrectal Transvaginal											
Transurethral Intravascular		ļ		<u> </u>						· · · · · · · · · · · · · · · · · · ·	
Peripheral vessel		P	P	P		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11, 14	
Laparoscopic										Note	
Musculo-skeletal Conventional		P	P	P		P	Р		BMDC	2,3,4,5,7,8,10, 11, 14	
Musculo-skeletal Superficial		Р	Р	P		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11, 14	
Other (specify)	1		İ								

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc.

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 53-Scape real-time 3D imagingNote 7B&W SieScape panoramic imagingNote 8Power SieScape panoramic imagingNote 10 Clarify VE vascular enhancement technologyNote 11Advanced Sieclear multi-view spatial compoundingNote 14eSie™ Touch elasticity imaging/FTI

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

()00

Ultrasound Indications for Use Form

510 (k) Number (if known):

K081148

Device Name:

14L5 Multi-D Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation									
Clinical Application	А	В	M	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)
Ophthalmic									,	
Fetal										
Abdominal				1						
Intraoperative Abdominal										•
Intraoperative Neurological										
Pediatric				1	<u> </u>					
Small Organ (Note 1)		N					<u>.</u>			Note 2,3,4,5,7,8,10, 11, 14
Neonatal Cephalic										
Adult Cephalic										
Cardiac										
Trans- esophageal										
Transrectal				-l	<u> </u>				+	
Transvaginal		<u> </u>			l					
Transurethral										
Intravascular Peripheral vessel	<u> </u>	1		+					<u> </u>	
13		<u> </u>					<u> </u>			
Laparoscopic Musculo-skeletal		<u> </u>	1	+						
Conventional										
Musculo-skeletal Superficial										
Other (specify)										

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc.

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 6 Cadence contrast agent imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

510 (k) Number (if known):

Ko81148

Device Name:

4P1 Phased Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application	А	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Ρ	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10	
Abdominal		P	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10	
Intraoperative Abdominal											
Intraoperative Neurological			Ň								
Pediatric			1	1							
Small Organ			[1							
Neonatal Cephalic											
Adult Cephalic		Р	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10	
Cardiac		Р	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,6,7,8,10	
Trans-											
esophageal											
Transrectal											
Transvaginal									<u> </u>		
Transurethral								L	ļ		
Intravascular						·			1	· · · · · · · · · · · · · · · · · · ·	
Peripheral vessel						<u> </u>	ļ	<u> </u>		<u></u>	
Laparoscopic											
Musculo-skeletal						1					
Conventional	<u> </u>	ļ				<u> </u>		<u> </u>			
Musculo-skeletal						1					
Superficial			-		-	+			-		
Other (specify)							<u> </u>	<u> </u>	<u> </u>		

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

- Note 4 Tissue Equalization Technology
- Note 5 3-Scape real-time 3D imaging
- Note 6 Cadence contrast agent imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number

510 (k) Number (if known):

K081148

Device Name:

Intended Use:

6C2 Curved Array Transducer for use with ACUSON S2000 ABVS Ultrasound System Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation										
Clinical Application	A	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Color Velocity Imaging	Combin ed (Specify)	Other (Specify)		
Ophthalmic												
Fetal		Ρ	Р	Ρ		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11		
Abdominal		Р	Р	Р		Р	P		BMDC	Note 2,3,4,5,7,8,10, 11		
Intraoperative Abdominal												
Intraoperative Neurological												
Pediatric		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11		
Small Organ												
Neonatal Cephalic												
Adult Cephalic			_									
Cardiac												
Trans- esophageal												
Transrectal												
Transvaginal								-				
Transurethral		<u> </u>										
Intravascular			. <u></u>	··								
Peripheral vessel		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11		
Laparoscopic												
Musculo-skeletal												
Conventional			<u> </u>									
Musculo-skeletal Superficial												
Other (specify)												

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE/DØ NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE) Prescription Use (Per 21 CFR 801.109) Diagnostic Ultrasound Indications for Use Form

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 581148 510(k) Number .

510 (k) Number (if known):

<041148

Device Name:

4C1 Curved Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application	A	в	М	PWD	CWD	Color Doppler	Amplitud e Doppler	Color Velocity Imaging	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Ρ	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11	
Abdominal		P	Р	Ρ		Р	Р		BMDC	Note2,3,4,5,6,7,8 , 10, 11,	
Intraoperative Abdominal											
Intraoperative Neurological											
Pediatric Small Organ											
Neonatal Cephalic											
Adult Cephalic Cardiac											
Trans- esophageal											
Transrectal Transvaginal					1						
Transurethral						-					
Intravascular Peripheral vessel											
Laparoscopic Musculo-skeletal		<u> </u>								· ·	
Conventional Musculo-skeletal		<u> </u>							<u> </u>		
Superficial Other (specify)		 			<u> </u>						

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 6 Cadence contrast agent imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Diagnostic Ultrasound Indications for Use Form

510 (k) Number (if known):

(Division Sign-Off) Division of Reproductive, Abdominal and **Radiological Devices** 681 510(k) Number

Device Name:

4V1 Phased Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation										
Clinical Application	А	В	м	PWD	CWD	Color Dopple r	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)		
Ophthalmic												
Fetal		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10		
Abdominal		Р	Р	Ρ		Р	Ρ		BMDC	Note 2,3,4,5,7,8,10,		
Intraoperative Abdominal												
Intraoperative Neurological												
Pediatric												
Small Organ Neonatal Cephalic					-							
Adult Cephalic Cardiac												
Trans- esophageal												
Transrectal Transvaginal												
Transurethral									· · · · · · · · · · · · · · · · · · ·			
Intravascular Peripheral vessel		<u> </u>			<u> </u>							
Laparoscopic Musculo-skeletal Conventional												
Musculo-skeletal Superficial Other (specify)												

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE) Prescription Use (Per 21 CFR 801.109)

lu

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number _____ Kのタル48

510 (k) Number (if known):

081148

Device Name:

10V4 Phased Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation										
Clinical Application	A	В	М	PWD	CWD	Color Doppler	Amplitud e Doppler	Color Velocity Imaging	Combin ed (Specify)	Other (Specify)		
Ophthalmic				:			-					
Fetal		Ρ	Ρ	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10		
Abdominal		Р	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10		
Intraoperative Abdominal												
Intraoperative Neurological												
Pediatric		Р	Р	Р	Р	Р	Р		BMDC	Note 2,3,4,5,7,8,10		
Small Organ												
Neonatal Cephalic		Р	Р	Р	P	Р	Р		BMDC	Note 2,3,4,5,7,8,10		
Adult Cephalic										٤		
Cardiac		Ρ	Ρ	Ρ	Р	P	P		BMDC	Note 3,4		
Trans- esophageal												
Transrectal		-										
Transvaginal		1						<u> </u>				
Transurethral			<u> </u>				· .					
Intravascular		ļ	 	<u> </u>				ļ	_			
Peripheral vessel		Р	Р	Р	Р	P	P		BMDC	Note 2,3,4,5,7,8,10		
Laparoscopic									<u> </u>			
Musculo-skeletal Conventional												
Musculo-skeletal Superficial												
Other (specify)	1	1		1	1	1-1						

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (QDE) Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and **Radiological Devices** 510(k) Number

510 (k) Number (if known):

K081148

Device Name:

14L5 SP Linear Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Indications For Use:

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

	Mode of Operation									
Clinical Application	A	в	М	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combine d (Specify)	Other (Specify)
Ophthalmic										
Fetal										
Abdominal										
Intraoperative (Note 9)		Ρ	Ρ	Р		Р	P		BMDC	Note 2,3,4,5,7,8,10
Intraoperative Neurological		Ρ	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11
Pediatric					[
Small Organ (Note 1)		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14
Neonatal Cephalic										
Adult Cephalic										
Cardiac				<u> </u>				1		
Transesophageal				1					· · · · ·	
Transrectal										······································
Transvaginal										
Transurethral	1			1	1					
Intravascular			[
Peripheral vessel		Р	Р	Р		Р	Р		BMDC	Note2,3,4,5,6 ,7,8,10, 11 <u>,</u> 14
Laparoscopic										
Musculo-skeletal Conventional		Р	P	P		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14
Musculo-skeletal Superficial										
Other (specify)										

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc.

- Note 2 Ensemble tissue harmonic imaging
- Note 3 SieClear multi-view spatial compounding
- Note 4 Tissue Equalization Technology
- Note 5 3-Scape real-time 3D imaging
- Note 6 Cadence contrast agent imaging
- Note 7 B&W SieScape panoramic imaging
- Note 8 Power SieScape panoramic imaging
- Note 9 For example: vascular, abdominal

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number ______KO8(148)

510 (k) Number (if known):

K081148

Device Name:

7CF2 Curved array mechanical 3D transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation									
Clinical Application	А	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Ρ	Ρ	Ρ.		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,13	
Abdominal		Ρ	Р	Р		Ρ	Р		BMDC	Note 2,3,4,5,7,8,10, 11, 13	
Intraoperative Abdominal				·····			· · · · •				
Intraoperative Neurological											
Pediatric											
Small Organ											
Neonatal Cephalic											
Adult Cephalic											
Cardiac											
Trans-											
esophageal											
Transrectal											
Transvaginal											
Transurethral			~·								
Intravascular											
Peripheral vessel				·							
Laparoscopic											
Musculo-skeletal											
Conventional Musculo-skeletal						*					
Superficial											
Other (specify)											
	L	L		L					1		

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

Note 13 STIC

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

(Division Slgn-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number

510 (k) Number (if known):

681148

Device Name:

9EVF4 Curved Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application	A	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)	
Ophthalmic											
Fetal		Р	Ρ	Р		P	Р		BMDC	Note 2,3,4,5,7 <u>,</u> 8, 10,11	
Abdominal											
Intraoperative Abdominal											
Intraoperative Neurological								,			
Pediatric										•	
Small Organ											
Neonatal Cephalic		P	Р	P		Р	P		BMDC	Note 2,3,4,5,7,8, 10,11	
Adult Cephalic											
Cardiac											
Trans- esophageal											
Transrectal		ļ									
Transvaginal		P	P	P		P	P		BMDC	Note 2,3,4,5,7,8, 10,11	
Transurethral											
Intravascular											
Peripheral vessel											
Laparoscopic											
Musculo-skeletal											
Conventional				ļ			 	ļ		· · · ·	
Musculo-skeletal											
Superficial	ļ	<u> </u>					 				
Other (specify)	-	<u> </u>	ļ	<u> </u>	<u> </u>	<u> </u>		L			

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

Note 11 Advanced Sieclear multi-view spatial compounding

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices KOBIHS 510(k) Number

K081148

510 (k) Number (if known):

Device Name:

V5Ms Multiplane TEE Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation										
Clinical Application	А	в	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Velocity	Combin ed (Specify)	Other (Specify)		
Ophthalmic												
Fetal												
Abdominal							<u> </u>					
Intraoperative												
Abdominal]	<u> </u>									
Intraoperative												
Neurological										······································		
Pediatric			L				l					
Small Organ		<u> </u>					· · · · · · · · · · · · · · · · · · ·					
Neonatal					1							
Cephalic												
Adult Cephalic					L							
Cardiac				ļ								
Trans-		P	р	P	P	Р	Р		BMDC			
esophageal		<u>'</u>	ļ	ļ ' .	<u> </u>	ļ	ļ					
Transrectal		<u> </u>			ļ					· · · · · · · · · · · · · · · · · · ·		
Transvaginal	<u> </u>	<u> </u>				<u> </u>				U . ¬		
Transurethral					ļ		ļ					
Intravascular					ļ	<u> </u>						
Peripheral vessel		<u> </u>		<u> </u>	<u> </u>	<u> </u>	ļ		·	L		
Laparoscopic	L	<u> </u>	L	<u> </u>	<u> </u>	<u> </u>	<u> </u>	ļ				
Musculo-skeletal												
Conventional	<u> </u>				<u> </u>		<u> </u>		ļ			
Musculo-skeletal						1						
Superficial		<u> </u>				L						
Other (specify)			1							l		

N = new indication; P = previously cleared by FDA K# 063803; E = added under Appendix E

Additional Comments: n/a

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

Veu

510 (k) Number (if known):

K081148

Device Name:

18L6HD Linear Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation									
Clinical Application	А	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Color Velocity Imaging	Combin ed (Specify)	Other (Specify)
Ophthalmic										
Fetal				1						
Abdominal										
Intraoperative Abdominal										
Intraoperative Neurological										
Pediatric										
Small Organ (Note 1)		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14
Neonatal Cephalic										
Adult Cephalic										
Cardiac										
Trans- esophageal										
Transrectal										
Transvaginal					ļ		ļ			
Transurethral										· · · · · · · · · · · · · · · · · · ·
Intravascular			,		ļ					
Peripheral vessel		Р	Р	P		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14
Laparoscopic										
Musculo-skeletal Conventional		Р	Р	Р		Р	Р		BMDC	Note 2,3,4,5,7,8,10, 11,14
Musculo-skeletal Superficial		Р	Р	P	 	Р	P		BMDC	Note 2,3,4,5,7,8,10, 11,14
Other (specify)										

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 1 For example: breast, testes, thyroid, penis, prostate, etc.

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

Note 4 Tissue Equalization Technology

Note 5 3-Scape real-time 3D imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology Note 11 Advanced Sieclear multi-view spatial compounding Note 14 eSie™ Touch elasticity imaging/FTI

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE) Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number

510 (k) Number (if known):

051148

Device Name:

8V3 Phased Array Transducer for use with ACUSON S2000 ABVS Ultrasound System

Intended Use:

Ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation											
Clinical Application	А	В	м	PWD	CWD	Color Doppler	Amplitud e Doppler	Color Velocity Imaging	Combin ed (Specify)	Other (Specify)			
Ophthalmic													
Fetal		Р	Ρ	Р	Р	Ρ	Р		BMDC	Note 2,3,4,5,7,8,10			
Abdominal										· · · · · · · · · · · · · · · · · · ·			
Intraoperative Abdominal													
Intraoperative Neurological													
Pediatric		Р	Р	Р	P	Р	Р		BMDC	Note 2,3,4,5,7,8,10			
Small Organ													
Neonatal Cephalic		Ρ	Р	Р	Ρ	Р	Р		BMDC	Note 2,3,4,5,7,8,10			
Adult Cephalic													
Cardiac		Р	P	Р	P	Р	Р		BMDC	Note 3,4,6			
Trans- esophageal													
Transrectal													
Transvaginal				_									
Transurethral													
Intravascular													
Peripheral vessel													
Laparoscopic										·			
Musculo-skeletal Conventional													
Musculo-skeletal Superficial				<u> </u>									
Other (specify) Neonatal Cardiac		Р	Р	Р	Р	Р	Р		BMDC	Note 3,4,6			

N = new indication; P = previously cleared by FDA K# 063085; E = added under Appendix E

Additional Comments:

Note 2 Ensemble tissue harmonic imaging

Note 3 SieClear multi-view spatial compounding

- Note 4 Tissue Equalization Technology
- Note 5 3-Scape real-time 3D imaging

Note 6 Cadence contrast agent imaging

Note 7 B&W SieScape panoramic imaging

Note 8 Power SieScape panoramic imaging

Note 10 Clarify VE vascular enhancement technology

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED) Concurrence of CDRH, Office of Device Evaluation (ODE)

Prescription Use (Per 21 CFR 801.109)

(Division Sign-Off) Division of Reproductive, Abdominal and Radiological Devices 510(k) Number ______K061148