§ 26.49

the extent possible. Such participation involves developing and reviewing documents developed by the GHTF and jointly determining whether they are applicable to the implementation of this subpart.

§26.49 Regulatory cooperation.

- (a) The parties and authorities shall inform and consult with one another, as permitted by law, of proposals to introduce new controls or to change existing technical regulations or inspection procedures and to provide the opportunity to comment on such proposals.
- (b) The parties shall notify each other in writing of any changes to appendix A of this subpart.

§ 26.50 Alert system and exchange of postmarket vigilance reports.

- (a) An alert system will be set up during the transition period and maintained thereafter by which the parties will notify each other when there is an immediate danger to public health. Elements of such a system will be described in an appendix F of this subpart. As part of that system, each party shall notify the other party of any confirmed problem reports, corrective actions, or recalls. These reports are regarded as part of ongoing investigations.
- (b) Contact points will be agreed between both parties to permit authorities to be made aware with the appropriate speed in case of quality defect, batch recalls, counterfeiting and other problems concerning quality, which could necessitate additional controls or suspension of the distribution of the product.
- APPENDIX A TO SUBPART B OF PART 26— RELEVANT LEGISLATION, REGULA-TIONS, AND PROCEDURES.
- 1. For the European Community (EC) the following legislation applies to $\S26.42(a)$ of this subpart:

[Copies of EC documents may be obtained from the European Document Research, 1100 17th St. NW., suite 301, Washington, DC 20036.]

- a. Council Directive 90/385/EEC of 20 June 1990 on active implantable medical devices
 - OJ No. L 189, 20.7. 1990, p. 17. Conformity assessment procedures.

Annex 2 (with the exception of section 4)

Annex 4 Annex 5

b. Council Directive 93/42/EEC of 14 June 1993 on Medical Devices OJ No. L 169,12.7.1993, p.1. Conformity assessment procedures.

Annex 2 (with the exception of section 4)

Annex 3 Annex 4

Annex 5

Annex 6

2. For the United States, the following legislation applies to §26.32(a):

[Copies of FDA documents may be obtained from the Government Printing Office, 1510 H St. NW., Washington, DC 20005. FDA documents may be viewed on FDA's Internet web site at http://www.fda.gov.]

- a. The Federal Food, Drug and Cosmetic Act, 21 U.S.C. 321 et seq.
- b. The Public Health Service Act, 42 U.S.C. $201 \ et \ seq.$
- c. Regulations of the United States Food and Drug Administration found at 21 CFR, in particular, Parts 800 to 1299.
- d. Medical Devices; Third Party Review of Selected Premarket Notifications; Pilot Program, 61 FR 14789-14796 (April 3, 1996).
- e. Draft Guidance Document on Accredited Persons Program, 63 FR 28392 (May 22, 1998). f. Draft Guidance for Staff, Industry and Third Parties, Third Party Programs under the Sectoral Annex on Medical Devices to the Agreement on Mutual Recognition Between the United States of America and the European Community (MRA), 63 FR 36240 (July 2, 1998).
- g. Guidance Document on Use of Standards, 63 FR 9561 (February 25, 1998).

APPENDIX B TO SUBPART B OF PART 26— SCOPE OF PRODUCT COVERAGE

1. Initial Coverage of the Transition Period

Upon entry into force of this subpart as described in §26.80 (it is understood that the date of entry into force will not occur prior to June 1, 1998, unless the parties decide otherwise), products qualifying for the transitional arrangements under this subpart include:

- a. All Class I products requiring premarket evaluations in the United States—see Table 1.
- b. Those Class II products listed in Table 2.

2. During the Transition Period

The parties will jointly identify additional product groups, including their related accessories, in line with their respective priorities as follows:

 a. Those for which review may be based primarily on written guidance which the parties will use their best efforts to prepare expeditiously; and

b. Those for which review may be based primarily on international standards, in order for the parties to gain the requisite

experience.

The corresponding additional product lists will be phased in on an annual basis. The parties may consult with industry and other interested parties in determining which products will be added.

- 3. Commencement of the Operational Period
- a. At the commencement of the operational period, product coverage shall extend to all Class I/II products covered during the transition period.
- b. FDA will expand the program to categories of Class II devices as is consistent with the results of the pilot, and with

Pt. 26, Subpt. B, App. B

FDA's ability to write guidance documents if the device pilot for the third party review of medical devices is successful. The MRA will cover to the maximum extent feasible all Class II devices listed in Table 3 for which FDA-accredited third party review is available in the United States.

4. Unless explicitly included by joint decision of the parties, this part does not cover any U.S. Class II-tier 3 or any Class III product under either system.

[The lists of medical devices included in these tables are subject to change as a result of the Food and Drug Administration Modernization Act of 1997.]

Table 1—Class I Products Requiring Premarket Evaluations in the United States, Included in Scope of Product Coverage at Beginning of Transition Period ¹

21 CFR Section No.	Regulation Name
	Product Code—Device Name
Anesthesiology Panel (21 CFR part 868)	
868.1910	Esophageal Stethoscope
868.5620	BZW—Stethoscope, Esophageal Breathing Mouthpiece BYP—Mouthpiece, Breathing
868.5640	Medicinal Nonventilatory Nebulizer (Atomizer) CCQ—Nebulizer, Medicinal, Nonventilatory (Atomizer)
868.5675	Rebreathing Device BYW—Device, Rebreathing
868.5700	Nonpowered Oxygen Tent FOG—Hood, Oxygen, Infant BYL—Tent, Oxygen
868.6810	Tracheobronchial Suction Catheter BSY—Catheters, Suction, Tracheobronchial
Cardiovascular Panel	
(None) Dental Panel (21 CFR part 872)	
872.3400	Karaya and Sodium Borate With or Without Acacia Denture
	Adhesive KOM—Adhesive, Denture, Acacia and Karaya With Sodium Borate
872.3700	Dental Mercury (U.S.P.) ELY—Mercury
872.4200	Dental Handpiece and Accessories EBW—Controller, Food, Handpiece and Cord EFB—Handpiece, Air-Powered, Dental EFA—Handpiece, Belt and/or Gear Driven, Dental EGS—Handpiece, Contra- and Right-Angle Attachment, Dental EKX—Handpiece, Direct Drive, AC-Powered
872.6640	EKY—Handpiece, Water-Powered Dental Operative Unit and Accessories
872.0040	EIA—Unit, Operative Dental
Ear, Nose, and Throat Panel (21 CFR part 874)	, ,
874.1070	Short Increment Sensitivity Index (SISI) Adapter ETR—Adapter, Short Increment Sensitivity Index (SISI)
874.1500	Gustometer ETM—Gustometer
874.1800	Air or Water Caloric Stimulator KHH—Stimulator, Caloric-Air ETP—Stimulator, Caloric-Water
874.1925	Toynbee Diagnostic Tube ETK—Tube, Toynbee Diagnostic
874.3300	Hearing Aid LRB—Face Plate Hearing-Aid
874.4100	ESD—Hearing-aid, Air-Conduction Epistaxis Balloon EMX—Balloon, Epistaxis

21 CFR Ch. I (4-1-11 Edition)

Pt. 26, Subpt. B, App. B

TABLE 1—CLASS I PRODUCTS REQUIRING PREMARKET EVALUATIONS IN THE UNITED STATES, INCLUDED IN SCOPE OF PRODUCT COVERAGE AT BEGINNING OF TRANSITION PERIOD 1—Continued

21 CFR Section No.	Regulation Name
	Product Code—Device Name
874.5300	ENT Examination and Treatment Unit
874.5550	ETF—Unit, Examining/Treatment, ENT Powered Nasal Irrigator
874.5840	KMA—Irrigator, Powered Nasal Antistammering Device
astroenterology—Urology Panel (21 CFR part 876)	KTH—Device, Anti-Stammering
876.5160	Urological Clamp for Males FHA—Clamp, Penile
876.5210	Enema Kit FCE—Kit, Enema, (for Cleaning Purpose)
876.5250	Urine Collector and Accessories FAQ—Bag, Urine Collection, Leg, for External Use
eneral Hospital Panel (21 CFR part 880)	
880.5270	Neonatal Eye Pad FOK—Pad, Neonatal Eye
880.5420	Pressure Infusor for an I.V. Bag
880.5680	KZD—Infusor, Pressure, for I.V. Bags Pediatric Position Holder
000.0000	FRP—Holder, Infant Position
880.6250	Patient Examination Glove
	LZB—Finger Cot FMC—Glove, Patient Examination
	LYY—Glove, Patient Examination, Latex LZA—Glove, Patient Examination, Poly LZC—Glove, Patient Examination, Speciality LYZ—Glove, Patient Examination, Vinyl
	LZA—Glove, Patient Examination, Poly
	LYZ—Glove, Patient Examination, Speciality LYZ—Glove Patient Examination, Vinyl
880.6375	Patient Lubricant
000.0700	KMJ—Lubricant, Patient
880.6760	Protective Restraint BRT—Restraint, Patient, Conductive
	FMQ—Restraint, Protective
eurology Panel (21 CFR part 882)	Atomiconante
882.1030	Ataxiagraph GWW—Ataxiagraph
882.1420	Electroencephalogram (EEG) Signal Spectrum Analyzer
992 4060	GWS—Analyzer, Spectrum, Electroencephalogram Signa Ventricular Cannula
882.4060	HCD—Cannula, Ventricular
882.4545	Shunt System Implantation Instrument GYK—Instrument, Shunt System Implantation
882.4650	Neurosurgical Suture Needle
992 4750	HAS—Needle, Neurosurgical Suture
882.4750	Skull Punch GXJ—Punch, Skull
bstetrics and Gynecology Panel (None)	
Ophthalmology Panel (21 CFR part 886)	
886.1780	Retinoscope HKM—Retinoscope, Battery-Powered
886.1940	Tonometer Sterilizer
886.4070	HKZ—Sterilizer, Tonometer Powered Corneal Burr
880.4070	HQS—Burr, Corneal, AC-Powered
	HOG—Burr, Corneal, Battery-Powered
	HRG—Engine, Trephine, Accessories, AC-Powered
	HFR—Engine, Trephine, Accessories, Battery-Powered HLD—Engine, Trephine, Accessories, Gas-Powered
886.4370	Keratome
	HNO—Keratome, AC-Powered
886.5850	HMY—Keratome, Battery-Powered Sunglasses (Nonprescription)
	HQY—Sunglasses (Nonprescription Including Photoset
rthopedic Panel (21 CFR part 888)	tive)
888.1500	Goniometer
	KQX—Goniometer, AC-Powered
888.4150	Calipers for Clinical Use

TABLE 1—CLASS I PRODUCTS REQUIRING PREMARKET EVALUATIONS IN THE UNITED STATES, INCLUDED IN SCOPE OF PRODUCT COVERAGE AT BEGINNING OF TRANSITION PERIOD 1—Continued

21 CFR Section No. Regulation Name	
	Product Code—Device Name
Physical Medicine Panel (21 CFR part 890)	
890.3850	Mechanical Wheelchair
	LBE—Stroller, Adaptive
	IOR—Wheelchair, Mechanical
890.5180	Manual Patient Rotation Bed
	INY—Bed, Patient Rotation, Manual
890.5710	Hot or Cold Disposable Pack
555.57.15	IMD—Pack, Hot or Cold, Disposable
adiology Panel (21 CFR part 892)	TWD T dok, Not of Cold, Dioposable
892.1100	Scintillation (Gamma) Camera
032.1100	IYX—Camera, Scintillation (Gamma)
892.1110	Positron Camera
892.1110	
000 4000	IZC—Camera, Positron
892.1300	Nuclear Rectilinear Scanner
	IYW—Scanner, Rectilinear, Nuclear
892.1320	Nuclear Uptake Probe
	IZD—Probe, Uptake, Nuclear
892.1330	Nuclear Whole Body Scanner
	JAM-Scanner, Whole Body, Nuclear
892.1410	Nuclear Electrocardiograph Synchronizer
	IVY—Synchronizer, Electrocardiograph, Nuclear
892.1890	Radiographic Film Illuminator
	IXC—Illuminator, Radiographic-Film
	JAG—Illuminator, Radiographic-Film, Explosion-Proof
892.1910	Radiographic Grid
692.1910	IXJ—Grid, Radiographic
000 4000	Deslie weeking between this work
892.1960	Radiographic Intensifying Screen
	EAM—Screen, Intensifying, Radiographic
892.1970	Radiographic ECG/Respirator Synchronizer
	IXO—Synchronizer, ECG/Respirator, Radiographic
892.5650	Manual Radionuclide Applicator System
	IWG—System, Applicator, Radionuclide, Manual
General and Plastic Surgery Panel (21 CFR part 878)	, , , , , , , , , , , , , , , , , , , ,
878.4200	Introduction/Drainage Catheter and Accessories
070.4200	KGZ—Accessories, Catheter
	GCE—Adaptor, Catheter
	FGY—Cannula, Injection GBA—Catheter, Balloon Type
	GBA—Catneter, Balloon Type
	GBZ—Catheter, Cholangiography
	GBQ—Catheter, Continuous Irrigation
	GBY—Catheter, Eustachian, General & Plastic Surgery
	JCY—Catheter, Infusion
	GBX—Catheter, Irrigation
	GBP—Catheter, Multiple Lumen
	GBO—Catheter, Nephrostomy, General & Plastic Surger
	CPN Cathotor Redictric Constal & Plantic Constant
	GBN—Catheter, Pediatric, General & Plastic Surgery GBW—Catheter, Peritoneal
	GBS—Catheter, Ventricular, General & Plastic Surgery
	GCD—Connector, Catheter
	GCC—Dilator, Catheter
	GCB-Needle, Catheter
878.4320	Removable Skin Clip
- =::===	FZQ—Clip, Removable (Skin)
878.4460	Surgeon's Gloves
O, O. TTOO	KGO—Surgeon's Gloves
070 4600	Nonpowored Single Potient Portable Susting Assessed
878.4680	Nonpowered, Single Patient, Portable Suction Apparatus
	GCY-Apparatus, Suction, Single Patient Use, Portal
	Nonpowered
878.4760	Removable Skin Staple
	GDT—Staple, Removable (Skin)
878.4820	AC-Powered, Battery-Powered, and Pneumatically Po
5.0.7020	ered Surgical Instrument Motors and Accessories/Atta
	ments
	GFG—Bit, Surgical
	GFA—Blade, Saw, General & Plastic Surgery
	DWH—Blade, Saw, Surgical, Cardiovascular
	BRZ—Board, Arm (With Cover)
	GFE—Brush, Dermabrasion

TABLE 1—CLASS I PRODUCTS REQUIRING PREMARKET EVALUATIONS IN THE UNITED STATES, INCLUDED IN SCOPE OF PRODUCT COVERAGE AT BEGINNING OF TRANSITION PERIOD 1—Continued

21 CFR Section No.	Regulation Name	
	Product Code—Device Name	
	KDG—Chisel (Osteotome)	
	GFD—Dermatome	
	GFC—Driver, Surgical, Pin	
	GFB—Head, Surgical, Hammer	
	GEY—Motor, Surgical Instrument, AC-Powered	
	GET-Motor, Surgical Instrument, Pneumatic Powered	
	DWI—Saw, Electrically Powered	
	KFK—Saw, Pneumatically Powered	
	HAB—Saw, Powered, and Accessories	
878.4960	Air or AC-Powered Operating Table and Air or AC-Powered Operating Chair & Accessories	
	GBB—Chair, Surgical, AC-Powered	
	FQO—Table, Operating-Room, AC-Powered	
	GDC—Table, Operating-Room, Electrical	
	FWW—Table, Operating-Room, Pneumatic	
	JEA—Table, Surgical with Orthopedic Accessories, AC-	
	Powered	
880.5090	Liquid Bandage	
	KMF—Bandage, Liquid	

¹Descriptive information on product codes, panel codes, and other medical device identifiers may be viewed on FDA's Internet Web Site at http://www.fda.gov/cdrh/prodcode.html.

Table 2—Class II Medical Devices Included in Scope of Product Coverage at Beginning of Transition Period (United States to develop guidance documents identifying U.S. requirements and European Community (EC) to identify standards needed to meet EC requirements) ¹

Panel	21 CFR Section No.	Regulation Name
		Product Code—Device Name
RA	892.1000	Magnetic Resonance Diagnostic Device MOS—COIL, Magnetic Resonance, Specialty LNH—System, Nuclear Magnetic Resonance Imaging LNI—System, Nuclear Magnetic Resonance Spectroscopic
Diagnostic Ultrasound:		
RA	892.1540	Nonfetal Ultrasonic Monitor JAF—Monitor, Ultrasonic, Nonfetal
RA	892.1550	Ultrasonic Pulsed Doppler Imaging System IYN—System, Imaging, Pulsed Doppler, Ultrasonic
RA	892.1560	Ultrasonic Pulsed Echo Imaging System IYO—System, Imaging, Pulsed Echo, Ultrasonic
RA	892.1570	Diagnostic Ultrasonic Transducer ITX—Transducer, Ultrasonic, Diagnostic
Diagnostic X-Ray Imaging Devices (except mammographic x-ray systems):		, A. J. Calabato, J. Calabato, J. Lagracia
RA	892.1600	Angiographic X-Ray System IZI—System, X-Ray, Angiographic
RA	892.1650	Image-Intensified Fluoroscopic X-Ray System MQB—Solid State X-Ray Imager (Flat Panel/Digital Imager) JAA—System, X-Ray, Fluoroscopic, Image-Intensified
RA	892.1680	Stationary X-Ray System KPR—System, X-Ray, Stationary
RA	892.1720	Mobile X-Ray System IZL—System, X-Ray, Mobile
RA	892.1740	Tomographic X-Ray System IZF—System, X-Ray, Tomographic
RA	892.1750	Computed Tomography X-Ray System JAK—System, X-Ray, Tomography, Computed
ECG-Related Devices:		orac Gystom, Array, romography, compated
CV	870.2340	Electrocardiograph DPS—Electrocardiograph MLC—Monitor. ST Segment
CV	870.2350	MILD—Monitor, ST Segment Electrocardiograph Lead Switching Adaptor DRW—Adaptor, Lead Switching, Electrocardiograph

TABLE 2—CLASS II MEDICAL DEVICES INCLUDED IN SCOPE OF PRODUCT COVERAGE AT BEGINNING OF TRANSITION PERIOD (UNITED STATES TO DEVELOP GUIDANCE DOCUMENTS IDENTIFYING U.S. REQUIREMENTS AND EUROPEAN COMMUNITY (EC) TO IDENTIFY STANDARDS NEEDED TO MEET EC REQUIREMENTS) 1—Continued

Panel	21 CFR Section No.	Regulation Name
		Product Code—Device Name
CV	870.2360	Electrocardiograph Electrode DRX—Electrode, Electrocardiograph
CV	870.2370	Electrocardiograph Surface Electrode Tester KRC—Tester, Electrode, Surface, Electrocardiographic
NE	882.1400	Electroencephalograph GWQ—Electroencephalograph
но	880.5725	Infusion Pump (external only) MRZ—Accessories, Pump, Infusion FRN—Pump, Infusion, Analytical Sampling MEB—Pump, Infusion, Elastomeric LZH—Pump, Infusion, Enteral MHD—Pump, Infusion, Gallstone Dissolution LZG—Pump, Infusion, Insulin
Ophthalmic Instru-		MEA—Pump, Infusion, PCA
ments:		
OP	886.1570	Ophthalmoscope HLI—Ophthalmoscope, AC-Powered HLJ—Ophthalmoscope, Battery-Powered
OP	886.1780	Retinoscope HKL—Retinoscope, AC-Powered
OP	886.1850	AC-Powered Slit-Lamp Biomicroscope HJO—Biomicroscope, Slit-Lamp, AC-Powered
OP	886.4150	Vitreous Aspiration and Cutting Instrument MMC—Dilator, Expansive Iris (Accessory) HQE—Instrument, Vitreous Aspiration and Cutting, AC-Powered HKP—Instrument, Vitreous Aspiration and Cutting, Battery-Powered MLZ—Vitrectomy, Instrument Cutter
OP	886.4670	Phacofragmentation System HQC—Unit, Phacofragmentation
SU	878.4580	Surgical Lamp HBI—Illuminator, Fiberoptic, Surgical Field FTF—Illuminator, Nonremote FTG—Illuminator, Remote HJE—Lamp, Fluorescein, AC-Powered FQP—Lamp, Operating-Room FTD—Lamp, Surgical GBC—Lamp, Surgical, Incandescent FTA—Light, Surgical, Accessories FSZ—Light, Surgical, Carrier FSY—Light, Surgical, Ceiling Mounted FSX—Light, Surgical, Connector FSW—Light, Surgical, Endoscopic FST—Light, Surgical, Fiberoptic FSS—Light, Surgical, Fibor Standing FSQ—Light, Surgical, Floor Standing FSQ—Light, Surgical, Instrument
NE	882.5890	Transcutaneous Electrical Nerve Stimulator for Pain Relief GZJ—Stimulator, Nerve, Transcutaneous, For Pain Relief Noninvasive Blood Pressure Measurement Devices:
CV	870.1120	Blood Pressure Cuff DXQ—Cuff, Blood-Pressure
CV	870.1130	Noninvasive Blood Pressure Measurement System (excep nonoscillometric) DXN—System, Measurement, Blood-Pressure, Noninvasive
НО	880.6880	Steam Sterilizer (greater than 2 cubic feet) FLE—Sterilizer, Steam
Clinical Thermometers:		
HO	880.2910	Clinical Electronic Thermometer (except tympanic or pacifier) FLL—Thermometer, Electronic, Clinical
AN	868.5630	Nebulizer CAF—Nebulizer (Direct Patient Interface)

Table 2—Class II Medical Devices Included in Scope of Product Coverage at Beginning of Transition Period (United States to develop guidance documents identifying U.S. requirements and European Community (EC) to identify standards needed to meet EC requirements) 1—Continued

Panel	21 CFR Section No.	Regulation Name
		Product Code—Device Name
Hypodermic Needles and Syringes (ex- cept antistick and self-destruct):		
НО	880.5570	Hypodermic Single Lumen Needle MMK—Container, Sharpes FMI—Needle, Hypodermic, Single Lumen MHC—Port, Intraosseous, Implanted
НО	880.5860	Piston Syringe FMF—Syringe, Piston
Selected Dental Materials:		· ···· · · · · · · · · · · · · · · · ·
DE	872.3060	Gold-Based Alloys and Precious Metal Alloys for Clinical Use EJT—Alloy, Gold Based, For Clinical Use EJS—Alloy, Precious Metal, For Clinical Use
DE	872.3200	Resin Tooth Bonding Agent KLE—Agent, Tooth Bonding, Resin
DE	872.3275	Dental Cement EMA—Cement, Dental EMB—Zinc Oxide Eugenol
DE	872.3660	Impression Material ELW—Material, Impression
DE	872.3690	Tooth Shade Resin Material EBF—Material, Tooth Shade, Resin
DE	872.3710	Base Metal Alloy EJH—Metal, Base
Latex Condoms:		
ОВ	884.5300	Condom HIS—Condom

¹Descriptive information on product codes, panel codes, and other medical device identifiers may be viewed on FDA's Internet Web Site at http://www.fda.gov/cdrh/prodcode.html.

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period ¹

Product Family	21 CFR Section No	Device Name	Tier
nesthesiology Panel			
Anesthesia Devices	868.5160	Gas machine for anesthesia or analgesia	2
	868.5270	Breathing system heater	2
	868.5440	Portable oxygen generator	2
	868.5450	Respiratory gas humidifier	2
	868.5630	Nebulizer	2
	868.5710	Electrically powered oxygen tent	2
	868.5880	Anesthetic vaporizer	2
Gas Analyser	868.1040	Powered Algesimeter	2
	868.1075	Argon gas analyzer	2
	868.1400	Carbon dioxide gas analyzer	2
	868.1430	Carbon monoxide gas ana- lyzer	2
	868.1500	Enflurane gas analyzer	2
	868.1620	Halothane gas analyzer	2
	868.1640	Helium gas analyzer	2
	868.1670	Neon gas analyzer	2
	868.1690	Nitrogen gas analyzer	2
	868.1700	Nitrous oxide gas analyzer	2
	868.1720	Oxygen gas analyzer	2
	868.1730	Oxygen uptake computer	2
Peripheral Nerve Stimulators	868.2775	Electrical peripheral nerve stimulator	2
Respiratory Monitoring	868.1750	Pressure plethysmograph	2
	868.1760	Volume plethysmograph	2
	868.1780	Inspiratory airway pressure meter	2

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family	21 CFR Section No	Device Name	Tier
	868.1800	Rhinoanemometer	2
	868.1840	Diagnostic spirometer	2
	868.1850	Monitoring spirometer	2
	868.1860	Peak-flow meter for	2
		spirometry	
	868.1880	Pulmonary-function data cal- culator	2
	868.1890	Predictive pulmonary-function value calculator	2
	868.1900	Diagnostic pulmonary-function interpretation calculator	2
	868.2025	Ultrasonic air embolism mon- itor	2
	868.2375	Breathing frequency monitor (except apnea detectors)	2
	868.2480	Cutaneous carbon dioxide (PcCO ₂) monitor	2
	868.2500	Cutaneous oxygen monitor (for an infant not under gas anesthesia)	2
	868.2550	Pneumotachomometer	2
	868.2600	Airway pressure monitor	2
	868.5665	Powered percussor	2
	868.5690	Incentive spirometer	2
Ventilator	868.5905	Noncontinuous ventilator (IPPB)	2
	868.5925	Powered emergency ventilator	2
	868.5935	External negative pressure ventilator	2
	868.5895	Continuous ventilator	2
	868.5955	Intermittent mandatory ventila- tion attachment	2
	868.6250	Portable air compressor	2
Cardiovascular Panel	000.0200	r ortable all compressor	_
Cardiovascular Diagnostic	870.1425	Programmable diagnostic computer	2
	870.1450	Densitometer	2
	870.2310	Apex cardiograph (vibrocardiograph)	2
	870.2320	Ballistocardiograph	2
	870.2340	Electrocardiograph	2
	870.2350	Electrocardiograph lead switching adaptor	1
	870.2360	Electrocardiograph electrode	2
	870.2370	Electrocardiograph surface electrode tester	2
	870.2400	Vectorcardiograph	1
	870.2450	Medical cathode-ray tube dis- play	1
	870.2675	Oscillometer	2
	870.2840	Apex cardiographic transducer	2
Cardiovascular Monitoring	870.2860	Heart sound transducer Valve, pressure relief, cardiopulmonary bypass	2
	870.1100	Blood pressure alarm	2
	870.1110	Blood pressure computer	2
	870.1110	Blood pressure cuff	2
	870.1130	Noninvasive blood pressure measurement system	2
	870.1140	Venous blood pressure ma- nometer	2
	870.1220	Electrode recording catheter or electrode recording probe	2
	870.1270	Intracavitary phonocatheter system	2
	870.1875	Stethoscope (electronic)	2
	870.2050	Biopotential amplifier and sig-	2
		nal conditioner	
	870.2060	Transducer signal amplifier and conditioner	2

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family	21 CFR Section No	Device Name	Tier
	870.2100	Cardiovascular blood flow- meter	2
	870.2120	Extravascular blood flow probe	2
	870.2300	Cardiac monitor (including cardiotachometer and rate alarm)	2
	870.2700	Oximeter	2
	870.2710	Ear oximeter	2
	870.2750	Impedance phlebograph	2
	870.2770	Impedance phiebograph	2
	870.2770 870.2780	Hydraulic, pneumatic, or pho-	2
	3.0.2100	toelectric plethysmographs	۷
	870.2850	Extravascular blood pressure	2
	0.0.2000	transducer	4
	870.2870	Catheter tip pressure trans-	2
	3.0.2070	ducer	_
	870.2880	Ultrasonic transducer	2
	870.2890	Vessel occlusion transducer	2
	870.2900	Patient transducer and elec-	2
	3.0.2300	trode cable (including con-	2
		nector)	
	870.2910	Radiofrequency physiological	2
	2.2.20.0	signal transmitter and re- ceiver	-
	870.2920	Telephone electrocardiograph transmitter and receiver	2
	870.4205	Cardiopulmonary bypass bub- ble detector	2
	870.4220	Cardiopulmonary bypass heart-lung machine console	2
	870.4240	Cardiovascular bypass heat exchanger	2
	870.4250	Cardiopulmonary bypass tem- perature controller	2
	870.4300	Cardiopulmonary bypass gas control unit	2
	870.4310	Cardiopulmonary bypass cor- onary pressure gauge	2
	870.4330	Cardiopulmonary bypass on- line blood gas monitor	2
	870.4340	Cardiopulmonary bypass level sensing monitor and/or con- trol	2
	870.4370	Roller-type cardiopulmonary bypass blood pump	2
	870.4380	Cardiopulmonary bypass pump speed control	2
	870.4410	Cardiopulmonary bypass in- line blood gas sensor	2
Cardiovascular Thera- peutic	870.5050	Patient care suction apparatus	2
	870.5900	Thermal regulation system	2
Defibrillator	870.5300	DC-defibrillator (including pad- dles)	2
	870.5325	Defibrillator tester	2
Echocardiograph	870.2330	Echocardiograph	2
Pacemaker & Acces-	870.1750	External programmable pace-	2
sories	870.3630	maker pulse generator Pacemaker generator function	2
	870.3640	analyzer Indirect pacemaker generator function analyzer	2
	870.3720	Pacemaker electrode function	2
Miscellaneous	870.1800	Withdrawal-infusion pump	2
	870.2800	Medical magnetic tape re- corder	2
	None	Batteries, rechargeable, class II devices	

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family	21 CFR Section No	Device Name	Tier
Dental Panel			
Dental Equipment	872.1720	Pulp tester	2
	872.1740	Caries detection device	2
	872.4120	Bone cutting instrument and	2
		accessories	
	872.4465	Gas-powered jet injector	2
	872.4475	Spring-powered jet injector	2
	872.4600	Intraoral ligature and wire lock	2
	872.4840	Rotary scaler	2
	872.4850	Ultrasonic scaler	2
	872.4920	Dental electrosurgical unit and accessories	2
	872.6070	Ultraviolet activator for polym- erization	2
	872.6350	Ultraviolet detector	2
Dental Material	872.3050	Amalgam alloy	2
	872.3060	Gold-based alloys and pre- cious metal alloys for clin- ical use	2
	872.3200	Resin tooth bonding agent	2
	872.3250	Calcium hydroxide cavity liner	2
	872.3260	Cavity varnish	2
	872.3275	Dental cement (other than zinc oxide-eugenol)	2
	872.3300	Hydrophilic resin coating for dentures	2
	872.3310	Coating material for resin fill- ings	2
	872.3590	Preformed plastic denture tooth	2
	872.3660	Impression material	2
	872.3690	Tooth shade resin material	2
	872.3710	Base metal alloy	2
	872.3750	Bracket adhesive resin and tooth conditioner	2
	872.3760	Denture relining, repairing, or rebasing resin	2
	872.3765	Pit and fissure sealant and conditioner	2
	872.3770	Temporary crown and bridge resin	2
	872.3820	Root canal filling resin (other than chloroform use)	2
	872.3920	Porcelain tooth	2
Dental X-ray	872.1800	Extraoral source x-ray system	2
,	872.1810	Intraoral source x-ray system	2
Dental Implants	872.4880	Intraosseous fixation screw or wire	2
	872.3890	Endodontic stabilizing splint	2
Orthodontic ar/Nose/Throat Panel	872.5470	Orthodontic plastic bracket	2
Diagnostic Equipment	874.1050	Audiometer	2
	874.1090	Auditory impedance tester	2
	874.1120	Electronic noise generator for audiometric testing	2
	874.1325	Electroglottograph	2
	874.1820	Surgical nerve stimulator/loca- tor	2
Hearing Aids	874.3300	Hearing aid (for bone-conduction)	2
	874.3310	Hearing aid calibrator and analysis system	2
	874.3320	Group hearing aid or group auditory trainer	2
	874.3330	Master hearing aid	2
Surgical Equipment	874.4250	Ear, nose, and throat electric or pneumatic surgical drill	1
	874.4490	Argon laser for otology, rhi- nology, and laryngology	2

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family	21 CFR Section No	Device Name	Tier
	874.4500	Ear, nose, and throat micro- surgical carbon dioxide laser	2
Gastroenterology/Urology Panel		14001	
Endoscope (including angioscopes, laparscopes, oph- thalmic endoscopes)	876.1500	Endoscope and accessories	2
	876.4300	Endoscopic electrosurgical unit and accessories	2
Gastroenterology	876.1725	Gastrointestinal motility moni- toring system	1
Hemodialysis	876.5600	Sorbent regenerated dialysate delivery system for hemo- dialysis	2
	876.5630	Peritoneal dialysis system and accessories	2
	876.5665	Water purification system for hemodialysis	2
	876.5820	Hemodialysis system and accessories	2
	876.5830	Hemodialyzer with disposable insert (kiil-type)	2
Lithotriptor	876.4500	Mechanical lithotriptor	2
Urology Equipment	876.1620	Urodynamics measurement system	2
	876.5320	Nonimplanted electrical con- tinence device	2
	876.5880	Isolated kidney perfusion and transport system and ac- cessories	2
General Hospital Panel Infusion Pumps and Systems	880.2420	Electronic monitor for gravity flow infusion systems	2
	880.2460	Electrically powered spinal fluid pressure monitor	2
	880.5430	Nonelectrically powered fluid injector	2
	880.5725	Infusion pump	2
Neonatal Incubators	880.5400	Neonatal incubator	2
	880.5410	Neonatal transport incubator	2
	880.5700	Neonatal phototherapy unit	2
Piston Syringes	880.5570	Hypodermic single lumen nee-	1
	880.5860	Piston syringe (except antistick)	1
	880.6920	Syringe needle introducer	2
Miscellaneous	880.2910	Clinical electronic thermom- eter	2
	880.2920 880.5100	Clinical mercury thermometer AC-powered adjustable hos-	2 1
	880.5500	pital bed AC-powered patient lift	2
	880.6880	Steam sterilizer (greater than 2 cubic feet)	2
leurology Panel			
	882.1020	Rigidity analyzer	2
	882.1610	Alpha monitor	2
Neuro-Diagnostic	882.1320	Cutaneous electrode	2 2
	882.1340	Nasopharyngeal electrode	2
	882.1350	Needle electrode	2
	882.1400	Electroencephalograph	2
	882.1460	Nystagmograph	2
	882.1480	Neurological endoscope	2
	882.1540	Galvanic skin response meas- urement device	2
	882.1550	Nerve conduction velocity measurement device	2
	882.1560	Skin potential measurement device	2

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

		ERIOD '—Continued	_
Product Family	21 CFR Section No	Device Name	Tier
	882.1570	Powered direct-contact tem- perature measurement de- vice	2
	882.1620	Intracranial pressure monitoring device	2
	882.1835	Physiological signal amplifier	2
	882.1845	Physiological signal condi- tioner	2
	882.1855	Electroencephalogram (EEG) telemetry system	2
	882.5050	Biofeedback device	2
Echoencephalography	882.1240	Echoencephalograph	2
RPG	882.4400	Radiofrequency lesion generator	2
Neuro Surgery	none 882.4305	Electrode, spinal epidural	2 2
	882.4305	Powered compound cranial drills, burrs, trephines, and their accessories	2
	882.4310	Powered simple cranial drills	2
		burrs, trephines, and their accessories	
	882.4360	Electric cranial drill motor	2
	882.4370	Pneumatic cranial drill motor	2
	882.4560	Stereotaxic instrument	2
	882.4725	Radiofrequency lesion probe	2
	882.4845 882.5500	Powered rongeur	2 2
Stimulators	882.1870	Lesion temperature monitor Evoked response electrical	2
	882.1880	stimulator Evoked response mechanical stimulator	2
	882.1890	Evoked response photic stim- ulator	2
	882.1900	Evoked response auditory stimulator	or
	882.1950	Tremor transducer	2
	882.5890	Transcutaneous electrical nerve stimulator for pain re- lief	2
Obstetrics/Gynecology Panel	004 4000	Toronto de al condesenso	0
Fetal Monitoring	884.1660	Transcervical endoscope (amnioscope) and acces- sories	2
	884.1690	Hysteroscope and acces-	2
	30.11000	sories (for performance standards)	_
	884.2225	Obstetric-gynecologic ultra- sonic imager	2
	884.2600	Fetal cardiac monitor	2
	884.2640	Fetal phonocardiographic monitor and accessories	2
	884.2660	Fetal ultrasonic monitor and accessories	2
	884.2675	Fetal scalp circular (spiral) electrode and applicator	1
	884.2700	Intrauterine pressure monitor and accessories	2
	884.2720	External uterine contraction monitor and accessories	2
	884.2740	Perinatal monitoring system and accessories	2
	884.2960	Obstetric ultrasonic transducer and accessories	2
Gynecological Surgery Equipment	884.1720	Gynecologic laparoscope and accessories	2
	884.4160 884.4550	Unipolar endoscopic coagu- lator-cutter and accessories	2
	884.4550 884.4120	Gynecologic surgical laser Gynecologic electrocautery	2
	884.4120 884.5300	and accessories Condom	2
	004.3300	Condon	4

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family		Povice Name	Tior
Product Family	21 CFR Section No	Device Name	Tier
Ophthalmic Implants	886.3320	Eye sphere implant	2
Contact Lens	886.1385	Polymethylmethacrylate (PMMA) diagnostic contact lens	2
	886.5916	Rigid gas permeable contact lens (daily wear only)	2
Diagnostic Equipment	886.1120	Opthalmic camera	1
	886.1220	Corneal electrode	1
	886.1250	Euthyscope (AC-powered)	1
	886.1360	Visual field laser instrument	1
	886.1510	Eye movement monitor	1
	886.1570 886.1630	Ophthalmoscope AC-powered photostimulator	1 1
	886.1640	Ophthalmic preamplifier	1
	886.1670	Ophthalmic isotope uptake probe	2
	886.1780	Retinoscope (AC-powered de- vice)	1
	886.1850	AC-powered slit lamp bio- microscope	1
	886.1930	Tonometer and accessories	2
	886.1945	Transilluminator (AC-powered device)	1
(Diamonti /2	886.3130	Ophthalmic conformer	2
(Diagnostic/Surgery Equipment)	886.4670	Phacofragmentation system	2
Ophthalmic Implants	886.3340 886.3800	Extraocular orbital implant Scleral shell	2
Surgical Equipment	880.5725	Infusion pump (performance standards)	2
	886.3100	Ophthalmic tantalum clip	2
	886.3300	Absorbable implant (scleral buckling method)	2
	886.4100	Radiofrequency electrosurgical cautery apparatus	2
	886.4115	Thermal cautery unit	2
	886.4150	Vitreous aspiration and cutting instrument	2
	886.4170	Cryophthalmic unit	2
	886.4250	Ophthalmic electrolysis unit (AC-powered device)	1
	886.4335	Operating headlamp (AC-pow- ered device)	1
	886.4390 886.4392	Ophthalmic laser Nd:YAG laser for posterior	2 2
	000.4392	capsulotomy	2
	886.4400	Electronic metal locator	1
	886.4440	AC-powered magnet	1
	886.4610	Ocular pressure applicator	2
	886.4690 886.4790	Ophthalmic photocoagulator	2 2
	886.5100	Ophthalmic sponge Ophthalmic beta radiation source	2
	none	Ophthalmoscopes, replace- ment batteries, hand-held	1
Orthopedic Panel			
Implants	888.3010	Bone fixation cerclage	2
	888.3020	Intramedullary fixation rod	2
	888.3030	Single/multiple component metallic bone fixation appli- ances and accessories	2
	888.3040	Smooth or threaded metallic bone fixation fastener	2
	888.3050	Spinal interlaminal fixation or- thosis	2
0 . 15	888.3060	Spinal intervertebral body fixa- tion orthosis	2
Surgical Equipment	888.1240	AC-powered dynamometer	2
	888.4580	Sonic surgical instrument and accessories/attachments	2

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1 —Continued

Product Family	21 CFR Section No	Device Name	Tier
	none	Accessories, fixation, spinal interlaminal	2
	none	Accessories, fixation, spinal intervertebral body	2
	none	Monitor, pressure, intracompartmental	1
	none	Orthosis, fixation, spinal intervertebral fusion	2
	none	Orthosis, spinal pedicle fixa- tion	
	none	System, cement removal ex- traction	1
Physical Medicine Panel Diagnostic Equipment or (Therapy) Therapeutic Equipment	890.1225	Chronaximeter	2
	890.1375	Diagnostic electromyograph	2
	890.1385	Diagnostic electromyograph needle electrode	2
	890.1450	Powered reflex hammer	2
	890.1850	Diagnostic muscle stimulator	2
or (Therapy)	890.5850	Powered muscle stimulator	2
Therapeutic Equipment	890.5100	Immersion hydrobath	2
	890.5110	Paraffin bath	2
	890.5500	Infrared lamp	2
	890.5720	Water circulating hot or cold pack	2
- diele en a Deur el	890.5740	Powered heating pad	2
ndiology Panel MRI	892.1000	Magnetic resonance diag- nostic device	2
Ultrasound Diagnostic	884.2660	Fetal ultrasonic monitor and accessories	2
	892.1540	Nonfetal ultrasonic monitor	
	892.1560	Ultrasonic pulsed echo imag- ing system	2
	892.1570	Diagnostic ultrasonic trans- ducer	2
	892.1550	Ultrasonic pulsed doppler im- aging system	
Angiographic	892.1600	Angiographic x-ray system	2
Diagnostic X-Ray	892.1610	Diagnostic x-ray beam-limiting device	2
	892.1620	Cine or spot fluorographic x- ray camera	2
	892.1630 892.1650	Electrostatic x-ray imaging system Image-intensified fluoroscopic	2
	892.1670	x-ray system Spot film device	2
	892.1680	Stationary x-ray system	2
	892.1710	Mammographic x-ray system	2
	892.1720	Mobile x-ray system	2
	892.1740	Tomographic x-ray system	1
	892.1820	Pneumoencephalographic chair	2
	892.1850	Radiographic film cassette	1
	892.1860	Radiographic film/cassette changer	1
	892.1870	Radiographic film/cassette changer programmer	2
	892.1900	Automatic radiographic film processor	2
	892.1980	Radiologic table	1
CT Scanner	892.1750	Computed tomography x-ray system	2
Radiation Therapy	892.5050	Medical charged-particle radi- ation therapy system	2
	892.5300	Medical neutron radiation therapy system	2

§ 26.60

Table 3—Medical Devices for Possible Inclusion in Scope of Product Coverage During Operational Period 1—Continued

Product Family	21 CFR Section No	Device Name	Tier
	892.5700	Remote controlled radio- nuclide applicator system	2
	892.5710	Radiation therapy beam-shap- ing block	2
	892.5730	Radionuclide brachytherapy source	2
	892.5750	Radionuclide radiation therapy system	2
	892.5770	Powered radiation therapy pa- tient support assembly	2
	892.5840	Radiation therapy simulation system	2
	892.5930	Therapeutic x-ray tube hous- ing assembly	1
Nuclear Medicine	892.1170	Bone densitometer	2
	892.1200	Emission computed tomog- raphy system	2
	892.1310	Nuclear tomography system	1
	892.1390	Radionuclide rebreathing system	2
eneral/Plastic Surgery Panel			
Surgical Lamps	878.4630	Ultraviolet lamp for dermato- logic disorders	2
	890.5500	Infrared lamp	2
	878.4580	Surgical lamp	2
Electrosurgical Cutting Equipment	878.4810	Laser surgical instrument for use in general and plastic surgery and in dermatology	2
	878.4400	Electrosurgical cutting and co- agulation device and acces- sories	2
Miscellaneous	878.4780	Powered suction pump	2

¹Descriptive information on product codes, panel codes, and other medical device identifiers may be viewed on FDA's Internet Web Site at http://www.fda.gov/cdrh/prodcode.html.

[63 FR 60141, Nov. 6, 1998; 64 FR 16348, Apr. 5, 1999]

APPENDICES C-F TO SUBPART B OF PART 26 [RESERVED]

Subpart C—"Framework" Provisions

§ 26.60 Definitions.

- (a) The following terms and definitions shall apply to this subpart only:
- (1) Designating Authority means a body with power to designate, monitor, suspend, remove suspension of, or withdraw conformity assessment bodies as specified under this part.
- (2) Designation means the identification by a designating authority of a conformity assessment body to perform conformity assessment procedures under this part.
- (3) Regulatory Authority means a government agency or entity that exercises a legal right to control the use or sale of products within a party's jurisdiction and may take enforcement ac-

tion to ensure that products marketed within its jurisdiction comply with legal requirements.

(b) Other terms concerning conformity assessment used in this part shall have the meaning given elsewhere in this part or in the definitions contained in "Guide 2: Standardization and Related Activities-General Vocabulary of the International Organization for Standardization (ISO) and the International Electrotechnical Commission (IEC)" (ISO/IEC Guide 2) (1996 edition), which is incorporated by reference in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. Copies are available from the International Organization for Standardization, 1, rue de Varembé, Case postale 56, CH-1211 Genève 20, Switzerland, or on the Internet at http://www.iso.ch or may be examined at the Food and Drug Administration's Medical Library, 5600 Fishers Lane, rm. 11B-40, Rockville, MD 20857,