regulation of a drug product under the act. Mr. Kokes was provided 30 days to file objections and request a hearing. Mr. Kokes did not request a hearing. His failure to request a hearing constitutes a waiver of his opportunity for a hearing and a waiver of any contentions concerning his debarment.

II. Findings and Order

Therefore, the Director, Center for Drug Evaluation and Research, under section 306(a)(2)(B) of the act, and under authority delegated to her (21 CFR 5.34), finds that Mr. Edwin Kokes has been convicted of a felony under Federal law for conduct relating to the regulation of a drug product under the act.

As a result of the foregoing finding, Mr. Edwin Kokes is permanently debarred from providing services in any capacity to a person with an approved or pending drug product application under sections 505, 512, or 802 of the act (21 U.S.C. 355, 360b, or 382) or under section 351 of the Public Health Service Act (42 U.S.C. 262)(see sections 306(c)(1)(B) and (c)(2)(A)(ii) and 201(dd) of the act (21 U.S.C. 321(dd))). Any person with an approved or pending drug product application who knowingly uses the services of Mr. Kokes, in any capacity, during his period of debarment, will be subject to civil money penalties (section 307(a)(6) of the act (21 U.S.C. 335b(a)(6))). If Mr. Kokes, during his period of debarment, provides services in any capacity to a person with an approved or pending drug product application, he will be subject to civil money penalties (section 307(a)(7) of the act). In addition, FDA will not accept or review any abbreviated new drug applications submitted by or with the assistance of Mr. Kokes during his period of debarment.

Any application by Mr. Kokes for termination of debarment under section 306(d)(4) of the act should be identified with Docket No. 01N–0539 and sent to the Dockets Management Branch (see **ADDRESSES**). All such submissions are to be filed in four copies. The public availability of information in these submissions is governed by 21 CFR 10.20(j). Publicly available submissions may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

Dated: March 28, 2003.

Steven K. Galson,

Deputy Director, Center for Drug Evaluation and Research.

[FR Doc. 03–10569 Filed 4–29–03; 8:45 am] BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 03N-0161]

Medical Devices; Reprocessed Single-Use Devices; Termination of Exemptions From Premarket Notification; Requirement for Submission of Validation Data

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a list (List I) of critical reprocessed single-use devices (SUDs) whose exemption from premarket submission is being terminated and for which validation data, as specified under the Medical Device User Fee and Modernization Act of 2002 (MDUFMA), is necessary in a premarket notification (510(k)); and a list (List II) of reprocessed SUDs that are currently subject to 510(k) requirements for which FDA has determined that validation data, as specified under MDUFMA, is necessary in a 510(k). FDA is requiring submission of these data to ensure that these reprocessed SUDs are substantially equivalent to predicate devices in accordance with MDUFMA.

DATES: These actions are effective April 30, 2003. Manufacturers of SUDs identified in List I whose exemption is being terminated must submit 510(k)s for these devices by July 30, 2004, or their devices may no longer be marketed. Manufacturers who already have clearance letters for SUDs identified in List II must submit validation data for these devices by January 30, 2004, or marketing of these devices must cease.

ADDRESSES: Submit written comments to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to *http:// www.fda.gov/dockets/ecomments*. Comments on Lists I and II should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

Barbara A. Zimmerman, Center for Devices and Radiological Health (HFZ– 410), Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850.

SUPPLEMENTARY INFORMATION:

I. Background

On October 26, 2002, MDUFMA (Pub. L. 107–250), amended the Federal Food, Drug, and Cosmetic Act (the act) by adding section 510(o) (21 U.S.C. 360(o)), which provided new regulatory requirements for reprocessed SUDs. According to this new provision, in order to ensure that reprocessed SUDs are substantially equivalent to predicate devices, 510(k)s for certain reprocessed SUDs identified by FDA must include validation data. These required validation data include cleaning and sterilization data, and functional performance data demonstrating that each SUD will remain substantially equivalent to its predicate device after the maximum number of times the device is reprocessed as intended by the person submitting the premarket notification.

Before enactment of the new law, a manufacturer of a reprocessed SUD was required to obtain premarket approval or premarket clearance for the device, unless the device was exempt from premarket submission requirements. Under MDUFMA, some previously exempt reprocessed SUDs will no longer be exempt from premarket notification requirements. Manufacturers of these identified devices will need to submit 510(k)s that include validation data to be specified by FDA. Reprocessors of certain SUDs that are currently subject to cleared 510(k)s also will need to submit the validation data specified by the agency.

In the near future, FDA will publish a guidance document providing more specific information about the types of validation data that should be submitted in premarket notification submissions for the reprocessed SUDs listed in this notice.

A. Definitions

Under section 302(b) of MDUFMA, a reprocessed SUD is defined as an "original device that has previously been used on a patient and has been subjected to additional processing and manufacturing for the purpose of an additional single use on a patient. The subsequent processing and manufacture of a reprocessed single-use device shall result in a device that is reprocessed within the meaning of this definition."

B. Reprocessed SUDs Exempt From Premarket Notification

Reprocessed SUDs are divided into three groups: (1) Critical, (2) semicritical, and (3) noncritical. The first two categories reflect definitions set forth in MDUFMA, and all three reflect a classification scheme recognized in the industry.¹ These categories of devices are defined as follows:

(1) A critical reprocessed SUD is intended to contact normally sterile tissue or body spaces during use.

(2) A semicritical reprocessed SUD is intended to contact intact mucous membranes and not penetrate normally sterile areas of the body.

(3) A noncritical reprocessed SUD is intended to make topical contact and not penetrate intact skin.

1. Requirements for Critical Reprocessed SUDs

MDUFMA requires FDA to review the critical reprocessed SUDs that are currently exempt from premarket notification requirements and determine which of these devices require premarket notification to ensure their substantial equivalence to predicate devices. By April 26, 2003, FDA must identify in a **Federal Register** notice those critical reprocessed SUDs whose exemption from premarket notification will be terminated. List I in this **Federal Register** notice implements this MDUFMA requirement.

In accordance with MDUFMA, manufacturers of the devices identified in List I must submit 510(k)s that include validation data regarding cleaning, sterilization, and functional performance, in addition to all the other required elements of a 510(k) identified in § 807.87 (21 CFR 807.87), within 15 months of publication of this notice or no longer market their device.

2. Requirements for Semicritical Reprocessed SUDs

MDUFMA also requires FDA to review the semicritical reprocessed SUDs that are currently exempt from premarket notification requirements and determine which of these devices require premarket notification to ensure their substantial equivalence to predicate devices. FDA must identify these devices in a notice published in the Federal Register by April 26, 2004. Manufacturers of devices identified at that time will be required to submit 510(k)s that include validation data regarding cleaning, sterilization, and functional performance in addition to all the other required elements of a 510(k) identified in § 807.87, within 15 months of publication of that notice or no longer market their device.

3. Requirements for Noncritical Reprocessed SUDs

MDUFMA does not require FDA to take any action under this section for noncritical SUDs that are exempt from premarket submission requirements.

C. Reprocessed SUDs Already Subject to Premarket Notification Requirements

MDUFMA also requires FDA to review the types of reprocessed SUDs already subject to premarket notification requirements and to identify which of these devices require the submission of validation data to ensure their substantial equivalence to predicate devices. FDA must publish a list of these devices in the **Federal Register** by April 26, 2003, and update the list as necessary. List II of this **Federal Register** notice implements this MDUFMA requirement. The devices on List II may be critical, semicritical, or noncritical reprocessed SUDs.

1. For devices identified in List II that have not yet been cleared through the 510(k) process, manufacturers must submit 510(k)s that include validation data regarding cleaning, sterilization, and functional performance, in addition to all the other required elements of a 510(k) identified in § 807.87, upon publication of this notice in order to market these devices. FDA will soon publish guidance to help submitters understand what types of validation data should be included in these 510(k)s.

2. For devices identified in List II that already have been cleared through the 510(k) process, manufacturers must submit validation data regarding cleaning, sterilization, and functional performance within nine months of publication of this notice or marketing must cease. FDA will soon publish guidance to explain how a 510(k) holder may submit the additional data now being required to support an earlier clearance.

II. FDA's Implementation of New Section 510(o) of the Act

FDA used a number of criteria to determine which device types should be included in the lists required by MDUFMA. As part of its consideration, FDA relied upon the Review Prioritization Scheme (RPS) it described in the February 2000 draft guidance document entitled "Reprocessing and Reuse of Single-Use Devices: Review Prioritization Scheme."² In the RPS guidance, FDA set forth factors that could be used to evaluate risk associated

with reprocessed SUDs. This approach assigned an overall risk to each SUD based on: (1) The risk of infection and (2) the risk of inadequate performance following reprocessing. Based on these risk factors, three categories of risk (high, moderate, and low) were developed. The designation of "high risk" was assigned to those devices that posed the greatest risk of infection and inadequate performance after reprocessing. In response to several comments about potential subjectivity of the RPS, FDA did not use the RPS approach when the agency finalized its enforcement priorities for reprocessed SUDs on August 14, 2000.

FDA has determined, however, that the RPS is an appropriate risk-based tool for developing the lists required by MDUFMA because the RPS identifies the devices that are likely to raise the most concerns about both infection transmission and inadequate performance following reprocessing. In formulating these lists, the agency also had the benefit of comments from stakeholders and an internal centerwide committee to evaluate the results of the RPS and ensure its consistency. In addition, there was a final review of all the devices on these lists by the Director of the Office of Device Evaluation. In this context, the agency believes these steps have adequately addressed concerns about the subjectivity of the RPS.

In addition to the previous criterion, FDA used one other criterion to identify those reprocessed SUDs that will be subject to the new requirements established by MDUFMA. The agency has included in these lists all reprocessed SUDs intended to come in contact with tissue at high risk of being infected with the causative agents of Creutzfeldt-Jakob Disease (CJD). These are generally devices intended for use in neurosurgery ophthalmology. This criterion was included in FDA's evaluation because insufficient scientific information exists at this time to establish standard methods to eliminate CJD infectious agents.

Therefore, in order to develop the two lists required by MDUFMA, FDA used the following process. First, the agency identified the types of SUDs that are being reprocessed. FDA did this by searching the 510(k) database for any 510(k)s that had been submitted for reprocessed SUDs and by asking original equipment manufacturers and reprocessors to provide information about types of devices that were being reprocessed. Second, FDA determined whether these devices are "critical," "semi-critical," or "non-critical". (These

¹ Spaulding, E.H., "The Role of Chemical Disinfection in the Prevention of Nonsocomial Infections," P.S. Brachman and T.C. Eickof (ed), Proceedings of International Conference on Nonsocomial Infections, 1970, American Hospital Association, Chicago, 1971:254–274.

² This draft guidance document is available on the CDRH Web site at *http//www.fda.gov/cdrh/ reuse/1156.pdf.*

definitions reflect the Spaulding ³ classification and are the same definitions FDA used earlier in developing its RPS.) FDA then applied the criteria described previously and "listed" any reprocessed SUD that was either "high" risk according to the RPS or intended to come in contact with tissue at high risk of being infected with the causative agents of CJD.

All devices identified in List I (previously exempt from 510(k)) have been determined to be critical reprocessed SUDs. In addition to being critical, they are either high risk according to the RPS or intended to come in contact with tissue at high risk of being infected with CJD. It should be noted that not all exempt devices that are critical have been listed. Critical reprocessed SUDs that are not listed in List I at this time may be reconsidered in subsequent updates of the list. The devices in List II (devices currently subject to 510(k) requirements that now will require the submission of validation data) are either high risk according to the RPS or intended to come in contact with tissue at high risk of being infected with the causative agents of CJD.

FDA has also provided a reference list in Attachment 1. To show how FDA evaluated the risk of a specific device, Attachment 1 includes the entire group of devices FDA considered when identifying the reprocessed SUDs in Lists I and II, and shows how FDA applied the criteria that determined whether the device would be identified on either of these lists.

In the Federal Register of February 4, 2003 (68 FR 5643), FDA invited interested persons to provide information and share views on the implementation of MDUFMA. The agency received several comments that identified specific reprocessed SUDs to be included in Lists I and II. The agency considered these recommendations while finalizing this document. Although FDA's lists do not include all the reprocessed SUDs that were recommended, the agency believes that those devices that pose the greatest risk of infection transmission and inadequate performance have been identified. The agency recognizes, however, that these lists may need to be re-evaluated and updated over time. Therefore, FDA will consider comments from the public on additional devices that should be included in the lists at any time. The agency also notes that MDUFMA permits FDA to request validation data for a device type that is subject to 510(k) clearance but not yet

included in List II. If this were to occur, FDA would ensure that manufacturers were aware of this change in the 510(k) submission requirements for that type of device by promptly updating the list.

Finally, FDA received one comment that suggested the agency's prior determinations about risk associated with reprocessed SUDs precluded FDA from now requiring 510(k)s for devices that were previously exempt or additional data for devices that were already cleared. FDA believes that this comment ignores the existence of MDUFMA's requirements. It is true that FDA had initially developed a regulatory approach for reprocessed SUDs that sought to treat those devices and original devices in a similar manner and that FDA had not required additional data to be submitted for certain reprocessed SUDs under that approach. However, through MDUFMA Congress clearly stated its intent to have the agency re-examine its policy with respect to reprocessed SUDs and legislated additional controls for those devices. FDA is committed to fulfilling its responsibilities under MDUFMA. The development and publication of these lists is part of the agency's implementation of these new statutory provisions.

LIST I.—CRITICAL REPROCESSED SINGLE-USE DEVICES PREVIOUSLY EXEMPT FROM PREMARKET NOTIFICATION REQUIREMENTS THAT WILL NOW REQUIRE 510(K)S WITH VALIDATION DATA

[To be submitted by July 26, 2004]

21 CFR section	Classification name	Product code for Non-reprocessed device	Product code for reprocessed device	Product code name for reprocessed device
872.3240	Dental bur	Diamond Coated	NME	Dental diamond coated bur.
872.4535	Dental diamond instrument	DZP	NLD	Dental diamond instrument.
872.4730	Dental injection needle	DZM	NMW	Dental needle.
874.4140	Ear, nose, and throat bur	Microdebrider	NLY	ENT high speed microdebrider.
874.4140	Ear, nose, and throat bur	Diamond Coated	NLZ	ENT diamond coated bur.
874.4420	Ear, nose, throat manual surgical * * *	KAB, KBG, KCI	NLB	Laryngeal, Sinus, Tracheal trocar.
878.4200	Introduction/drainage catheter and accessories.	GCB	NMT	Catheter needle.
878.4800	Manual surgical instrument	MJG	NNA	Percutaneous biopsy device.
878.4800	Manual surgical instrument	FHR	NMU	Gastro-Urology needle.
878.4800	Manual surgical instrument for * * *	DWO	NLK	Cardiovascular biopsy needle.
878.4800	Manual surgical instrument for * * *	GAA	NNC	Aspiration and injection.
882.4190	Forming/cutting clip instrument	HBS	NMN	Forming/cutting clip instrument.
884.1730	Laparoscopic insufflator * * *	HIF	NMI	Laparoscopic insufflator and accessories.
884.4530	OB/GYN specialized manual instrument	HFB	NMG	Gynecological biopsy forceps.
886.4350	Manual ophthalmic surgical instrument	HNN	NLA	Ophthalmic knife.

Infections," P. S. Brachman and T. C. Eickof (ed), Proceedings of International Conference on

³ Spaulding, E. H., "The Role of Chemical Disinfection in the Prevention of Nonsocomial

Nonsocomial Infections, 1970, American Hospital Association, Chicago, 1971:254–274.

LIST II.—REPROCESSED SINGLE-USE DEVICES SUBJECT TO PREMARKET NOTIFICATION REQUIREMENTS THAT WILL NOW REQUIRE THE SUBMISSION OF VALIDATION DATA¹

[Manufacturers who already have 510(k) clearance for these devices must submit validation data by January 26, 2004. Any new 510(k) will require validation data upon publication of this list.]

21 CFR section	Classification name	Product code for non-reprocessed device	Product code for reprocessed device	Product code name for reprocessed device
Unclassified Unclassified	Oocyte aspiration needles Percutaneous transluminal	MHK LIT	NMO NMM	Oocyte aspiration needles. Transluminal peripheral angioplasty catheter.
	angioplasty catheter angioplasty catheter.			· · · · · · · · · · · · · · · · · · ·
Unclassified 868.5150	Ultrasonic surgical instrument Anesthesia conduction needle	LFL BSP	NLQ NNH	Ultrasonic scalpel. Anesthetic conduction needle (with or without in- troducer).
868.5150	Anesthesia conduction needle	MIA	NMR	Short term spinal needle.
868.5730	Tracheal tube	BTR	NMA	Tracheal tube (with or without connector).
868.5905	Noncontinuous (IPPB)	BZD	NMC	Noncontinuous ventilator (respirator) mask.
870.1200 870.1220		DQO DRF	NLI NLH	Angiography catheter. Electrode recording catheter.
870.1220		MTD	NLG	Intracardiac mapping catheter.
870.1230		DQE	NMB	Fiberoptic oximeter catheter.
870.1280		DRA	NKS	Steerable catheter.
870.1290		DXX	NKR	Steerable catheter control system.
870.1330 870.1390		DQX DRC	NKQ NMK	Catheter guide wire.
870.1650	Trocar Angiographic injector and syringe	DXT	NKT	Cardiovascular trocar. Angiographic injector and syringe.
870.1670		DQF	NKW	Injector for actuator syringe.
870.2700		MUD	NMD	Tissue saturation oximeter.
870.2700	Oximeter	DQA	NLF	Oximeter.
870.3535	Intra-aortic balloon and control system.	DSP	NKO	Intra-aortic balloon and control system.
870.4450		DXC	NMF	Vascular clamp.
870.4885		DWQ	NLJ	External vein stripper.
872.5470 874.4680		DYW BWH	NLC NLE	Orthodontic plastic bracket. Bronchoscope (nonrigid) biopsy forceps.
	and accessories.			
876.1075		FCG	NMX	G–U biopsy needle and needle set.
876.1075	Gastroenterology-urology biopsy instrument.	KNW	NLS	Biopsy instrument.
876.1500		FBK, FHP	NMY	Endoscopic needle.
876.1500		MPÁ	NKZ	Endoilluminator.
876.1500		GCJ	NLM	General and plastic surgery laparoscope.
876.1500		FHO	NLX	Spring-loaded Pneumoperitoneum Needle.
876.4300	and accessories.		NLW	Active urological electrosurgical electrode.
876.4300		FEH	NLV	Flexible suction coagulator electrode.
876.4300	and accessories.	KGE	NLU	Electric biopsy forceps.
876.4300	Endoscopic electrosurgical unit and accessories.	FDI	NLT	Flexible snare.
876.4300	and accessories.	KNS	NLR	Endoscopic (with or without accessories) Electrosurgical unit.
876.5010	Biliary catheter and accessories	FGE	NML	Biliary catheter.
876.5540	Blood access device and accessories.	LBW	NNF	Single needle dialysis set (co-axial flow).
876.5540	Blood access device and accessories.	FIE	NNE	Fistula needle.
876.5820	Hemodialysis systems and accessories.	FIF	NNG	Single needle dialysis set with and accessories uni-directional pump.
878.4300	Implantable clip	FZP	NMJ	Implantable clip.
878.4750		GDW	NLL	Implantable staple.
880.5570		FMI	NKK	Hypodermic single lumen needle.
880.5860 882.4300	, , ,	FMF HBG	NKN NLO	Piston syringe. (Manual) drills, burrs, burrs, trephines and acces-
882.4305	trephines and accessories.	HBG	NLO	sories. (Powered, compound) drills, burrs, trephines and
882.4310	burrs, trephines	HBE	NLN	accessories. (Simple, powered) drills, burrs, trephines and ac-
	burrs, trephines.			cessories.
884.1720	cessories.	HET	NMH	Gynecologic laparoscope (and accessories).
004.0400				unanated reproduction poodlo
884.6100	Assisted reproduction needle Keratome	MQE HMY, HNO	NNB NKY	Assisted reproduction needle. Keratome blade.

LIST II.—REPROCESSED SINGLE-USE DEVICES SUBJECT TO PREMARKET NOTIFICATION REQUIREMENTS THAT WILL NOW REQUIRE THE SUBMISSION OF VALIDATION DATA ¹—Continued

[Manufacturers who already have 510(k) clearance for these devices must submit validation data by January 26, 2004. Any new 510(k) will require validation data upon publication of this list.]

21 CFR section	Classification na	ame Product code fo non-reprocesse device		Product code name for reprocessed device
892.5730	Radionuclide bra source.	achytherapy IWF	NMP	Isotope needle.

¹Hemodialyzers have been excluded from this list because the reuse of hemodialyzers is addressed in "Guidance for Hemodialyzer Reuse Labeling" (final draft issued on October 6, 1995).

III. Comments

You may submit written or electronic comments on these lists to the Dockets Management Branch (see **ADDRESSES**). You may submit a single copy of an electronic comment to *http:// www.fda.gov/dockets/ecomments*. You should submit two copies of any mailed comments but individuals may submit one copy. You should identify your comment with the docket number found in brackets in the heading of this document. You may see any comments FDA receives in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

ATTACHMENT 1.—LIST OF SUDS KNOWN TO BE REPROCESSED OR CONSIDERED FOR REPROCESSING

	Medical specialty	Device type	Regulation No.	Class	Product code	Risk 1,2,3,3*	Critical/semi- critical/non- critical	Premarket exempt
1 2	Cardio Cardio	Cardiopulmonary Bypass Marker Percutaneous & Operative Transluminal Coronary	Unclassified post amend- ment	ш	MAB LOX	1 3	C C	N N
3	Cardio	Angioplasty Catheter (PCTA). Percutaneous Ablation Electrode	Post	ш	LPB	3	с	N
4	Cardio	Peripheral Transluminal	amendment Unclassified		LIT	3	с	N
5	Cardio	Angioplasty (PTA) Catheter. Blood-Pressure Cuff	870.1120	п	DXQ	1	N	N
6	Cardio	Angiography Catheter	870.1200		DQO	3	C	N
-	Cardio		870.1200		DQO	3	C	N
7	Cardio	Electrode Recording Catheter	870.1220		MTD	3	C	N
8 9	Cardio	High-Density Array Catheter Fiberoptic Oximeter Catheter	870.1220		DQE	3	c	N
	Cardio	Steerable Catheter	870.1230		DQE	3	C	N
10 11	Cardio				DXX	3	C	N
12	Cardio	Steerable Catheter Control System	870.1290 870.1330		DAA DQX	3	C	N
12		Guide Wire	870.1390		DQA	3	C	N
13	Cardio	Angiographic Needle	870.1390		DRC	3	C	N
14	Cardio	Syringes	870.1650		DXT	3	C	N
16	Cardio	Injector Type Syringe Actuator	870.1670		DQF	3	c	N
17	Cardio	Oximeter	870.2700		DQA	3	N	N
18	Cardio	Tissue Saturation Oximeter	870.2700		MUD	3	C	N
10	Cardio	Intra-Aortic Balloon System	870.3535		DSP	3	c	N
20	Cardio	Vascular Clamp	870.4450		DXC	3	c	N
20	Cardio	Device, Stabilizer, Heart	870.4500		MWS	2	c	Y
22	Cardio	External Vein Stripper	870.4885		DWQ	3	c	Ň
23	Cardio	Compressible Limb Sleeve	870.5800		JOW	1	N	N
24	Dental	Bur	872.3240	lï	EJL	1	C	Y
25	Dental	Diamond Coated Bur	872.3240		EJL	3	c	Ý
26	Dental	Diamond Instrument	872.4535	li	DZP	3	c	Ý
20	Dental	AC-Powered Bone Saw	872.4120		DZH	2	c	Ň
28	Dental	Manual Bone Drill and Wire Driver	872.4120		DZJ	2	c	N
29	Dental	Powered Bone Drill	872.4120	ii ii	DZI	2	C	N
30	Dental	Intraoral Drill	872.4130	lï	DZA	1	č	Y
31	Dental	Injection Needle	872.4730	li	DZM	3	č	Ý
32	Dental	Metal Orthodontic Bracket	872.5410	li	EJF	3	S	Ŷ
33	Dental	Plastic Orthodontic Bracket	872.5470		DYW	3	S	N
34	ENT	Bur	874.4140	lï	EQJ	1	C	Y
35	ENT	Diamond Coated Bur	874.4140	li	EQJ	3	č	Ý
36	ENT	Microdebrider	874.4140	li	EQJ	3	C	Ý
37	ENT	Microsurgical Argon Fiber Optic Laser Cable, For Uses Other Than Otology, Including Laryn- gology & General Use In Otolarngology.	874.4490	II	LMS	1	S	Ň
38	ENT	Microsurgical Argon Fiber Optic Laser Cable, For Use In Otology.	874.4490	П	LXR	1	S	N

ATTACHMENT 1.-LIST OF SUDS KNOWN TO BE REPROCESSED OR CONSIDERED FOR REPROCESSING-Continued

	Medical specialty	Device type	Regulation No.	Class	Product code	Risk 1,2,3,3*	Critical/semi- critical/non- critical	Premarket exempt
39	ENT	Microsurgical Carbon-Dioxide Fiber Optic Laser Cable.	874.4500	П	EWG	1	S	N
40	ENT	Bronchoscope Biopsy Forceps (Non-Rigid).	874.4680	П	BWH	3	S	N
41	ENT	Bronchoscope Biopsy Forceps (Rigid).	874.4680	11	JEK	1	S	N
42	Gastro/Urol- ogy.	Biopsy Forceps Cover	876.1075	1	FFF	1	S	Y
43	Gastro/Urol- ogy.	Biopsy Instrument	876.1075	II	KNW	3	S	N
44	Gastro/Urol- ogy.	Biopsy Needle Set	876.1075	II	FCG	3	S	N
45	Gastro/Urol- ogy.	Biopsy Punch	876.1075	II	FCI	2	S	N
46	Gastro/Urol- ogy.	Mechanical Biopsy Instrument	876.1075	П	FCF	2	S	N
47	Gastro/Urol- ogy.	Non-Electric Biopsy Forceps	876.1075	1	FCL	3	S	Y
48	Gastro/Urol- ogy.	Cytology Brush For Endoscope	876.1500	П	FDX	2	S	N
49	Gastro/Urol- ogy.	Endoscope Accessories	876.1500	II	KOG	2	S	N
50	Gastro/Urol- ogy.	Extraction Balloons/Baskets	876.1500	II	KOG	2	S	N
51	Gastro/Urol- ogy.	Endoscopic Needle	876.1500	II	FBK	3	с	N
52	Gastro/Urol- ogy.	Simple Pneumoperitoneum Needle	876.1500	Ш	FHP	3	С	N
53	Gastro/Urol- ogy.	Spring Loaded Pneumoperitoneum Needle.	876.1500	П	FHO	3	С	N
54	Gastro/Urol- ogy.	Active Electrosurgical Electrode	876.4300	II	FAS	3	S	N
55	Gastro/Urol- ogy.	Biliary Sphincterotomes	876.5010, 876.1500	П	FGE	3	S	N
56	Gastro/Urol- ogy.	Electric Biopsy Forceps	876.4300	П	KGE	3	S	N
57	Gastro/Urol- ogy.	Electrosurgical Endoscopic Unit (With Or Without Accessories).	876.4300	П	KNS	3	S	N
58	Gastro/Urol- ogy.	Flexible Snare	876.4300	II	FDI	3	S	N
59	Gastro/Urol- ogy.	Flexible Suction Coagulator Elec- trode.	876.4300	II	FEH	3	S	N
60	Gastro/Urol- ogy.	Flexible Stone Dislodger	876.4680	II	FGO	3	S	Y
61	Gastro/Urol- ogy.	Metal Stone Dislodger	876.4680	II	FFL	3	S	Y
62	Gastro/Urol- ogy.	Needle Holder	876.4730	1	FHQ	1	с	Y
63	Gastro/Urol- ogy.	Non-Electrical Snare	876.4730	I	FGX	1	S	Y
64	Gastro/Urol- ogy.	Urological Catheter	876.5130	II	KOD	2	S	N
65	Gastro/Urol- ogy.	Single Needle Dialysis Set	876.5540	П	LBW, FIE	3	С	N
66	Gastro/Urol- ogy.	Hemodialysis Blood Circuit Accessories.	876.5820	П	кос	2	S	N
67	Gastro/Urol- ogy.	Single Needle Dialysis Set	876.5820	П	FIF	3	С	N
68 69	GE/U General	Hemorrhoidal Ligator Implanted, Programmable Infusion	876.4400 Post-amend-	11 111	FHN LKK	2	C C	N N
70	Hospital General	Pump. Needle Destruction Device	ment Post-amend-		MTV	1	N	N
70	Hospital General	Non-Powered Flotation Therapy	ment 880.5150	1	IKY	2	N	Y
72	Hospital General	Mattress. Non AC-Powered Patient Lift	880.5510		FSA	2	N	Y
72	Hospital General	Alternating Pressure Air Flotation	880.5550		FNM	1	N	Y
10	Hospital	Mattress.	000.0000					'

ATTACHMENT 1.—LIST OF SUDS KNOWN TO BE REPROCESSED OR CONSIDERED FOR REPROCESSING—Continued

	Medical specialty	Device type	Regulation No.	Class	Product code	Risk 1,2,3,3*	Critical/semi- critical/non- critical	Premarket exempt
	General Hospital	Temperature Regulated Water Mattress.	880.5560	I	FOH	2	N	Y
75	General Hospital	Hypodermic Single Lumen Needle	880.5570	П	FMI	3	С	N
76	General Hospital	Piston Syringe	880.5860	П	FMF	3	С	N
77	General Hospital	Mattress Cover (Medical Purposes).	880.6190	I	FMW	2	N	Y
78	General Hospital	Disposable Medical Scissors	880.6820	I	JOK	1	N	Y
	General Hospital	Irrigating Syringe	880.6960	I	KYZ, KYY	1	С	Y
	Infection Control	Surgical Gowns	878.4040	II	FYA	1	С	N
	Lab	Blood Lancet	878.4800	1	FMK	1	С	Y
	Neuro	Clip Forming/Cutting Instrument,	882.4190		HBS	3*	C	Y
	Neuro	Drills, Burrs, Trephines & Acces- sories (Manual).	882.4300		HBG	3*	С	N
	Neuro	Drills, Burrs, Trephines & Acces- sories (Compound, Powered). Drills, Burrs, Trephines & Acces-	882.4305 882.4310	11	HBF	3* 3*	c c	N
	Neuro OB/GYN	sories (Simple, Powered). Oocyte aspiration needle	Unclassified		MHK	3	c	N
	OB/GYN	Laparoscope Accessories	884.1720	ï	HET	2	c	Y
	OB/GYN	Laparoscope Accessories	884.1720	II.	HET	3	c	Ň
	OB/GYN	Laparoscopic Dissectors	884.1720	1	HET	2	C	Y
90	OB/GYN	Laparoscopic Graspers	884.1720	1	HET	2	C	Y
	OB/GYN	Laparoscopic Scissors	884.1720	1	HET	2	C	Y
	OB/GYN	Insufflator Accessories (tubing, Verres needle, kits).	884.1730	II	HIF	3	С	Y
	OB/GYN OB/GYN	Laparoscopic Insufflator Endoscopic Electrocautery And	884.1730 884.4100		HIF HIM	2	N N	N N
		Accessories.						
	OB/GYN	Gynecologic Electrocautery (And Accessories).	884.4120		HGI	2	N	N
	OB/GYN	Endoscopic Bipolar Coagulator- Cutter (And Accessories).	884.4150		HIN	2	N	N
	OB/GYN	Culdoscopic Coagulator (And Accessories).	884.4160		HFI	2	N	N
	OB/GYN	Endoscopic Unipolar Coagulator- Cutter (And Accessories).	884.4160		KNF	2	N	N
	OB/GYN	Hysteroscopic Coagulator (And Accessories).	884.4160		HFH	2	N	N
	OB/GYN	Unipolar Laparoscopic Coagulator (And Accessories).	884.4160		HFG	2	N	N
	OB/GYN	Episiotomy Scissors	884.4520		HDK	1	C	Y Y
	OB/GYN OB/GYN	Umbilical Scissors Biopsy Forceps	884.4520 884.4530		HDJ HFB	1	C C	Y
	OB/GYN OB/GYN	Assisted reproduction needle	884.6100		MQE	3	c	N
	Ophthalmic	Endoilluminator	876.1500	ii	MPA	3*	c	N
	Ophthalmic	Surgical Drapes	878.4370	ü	KKX	2	č	N
	Ophthalmic	Ophthalmic Knife	886.4350	i i	HNN	3	Ċ	Y
	Ophthalmic	Keratome Blade	886.4370	Inot ex- empt	HMY, HNO	3	C	N
	Ophthalmic Ophthalmic	Phacoemulsification Needle Phacoemulsification/ Phacofragmentation Fluidic.	886.4670 886.4670	II II	HQC MUS	3 2	C C	N N
111	Ophthalmic	Phacofragmentation Unit	886.4670	П	HQC	1	N	N
112	Ortho	Saw Blades	878.4820	1	GFA, DWH, GEY,	1	С	Y
113	Ortho	Surgical Drills	878.4820	I	GET GEY,	1	с	Y
114	Ortho	Arthroscope accessories	888.1100	П	GET HRX	2	с	Y
	Ortho	Bone Tap	888.4540	lï	HWX	2	c	Y
	Ortho	Burr	888.4540	i	HTT	1	c	Ý
	Ortho	Carpal Tunnel Blade	888.4540	1	LXH	2	c	Ý
118	Ortho	Countersink	888.4540	I	HWW	1	С	Y
119	Ortho	Drill Bit	888.4540	1	HTW	1	C	Υ

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ATTACHMENT 1.-LIST OF SUDS KNOWN TO BE REPROCESSED OR CONSIDERED FOR REPROCESSING-Continued

121 Ortti 122 Ortti 123 Ortti 124 Ortti 125 Ortti 126 Ortti 127 Ortti 128 Ortti 129 Ortti 130 Phy 131 Phy 133 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur Sur 137 Rac 138 Res	edicine	Knife Manual Surgical Instrument Needle Holder Reamer Scissors Staple Driver Trephine Flexible Reamers/Drills External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist. External Limb Component, Shoul-	888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3040 888.3030 Unclassified 890.1175 890.3420		HTS LXH HXK HTO HTX HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD ISL	1 1 1 1 1 1 1 2 1 1 2	CCCCCCCC N N N	Y Y Y Y Y Y Y Y Y Y
122 Ortl 123 Ortl 124 Ortl 125 Ortl 126 Ortl 127 Ortl 128 Ortl 129 Ortl 130 Phy 131 Phy 132 Phy 133 Phy 134 Phy 135 Phy 136 Pha 137 Rac 138 Res	rtho rtho rtho rtho rtho rtho rtho nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	Needle Holder Reamer	888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HXK HTO HTX HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 1 1 1 1 2 1 1	CCCCCCC N	Y Y Y Y Y Y N N
123 Ortl 124 Ortl 125 Ortl 126 Ortl 127 Ortl 128 Ortl 129 Ortl 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	rtho rtho rtho rtho rtho rtho nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	Needle Holder Reamer	888.4540 888.4540 888.4540 888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HXK HTO HTX HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 1 1 1 1 2 1 1	CCCCCCC N	Y Y Y Y Y Y N N
123 Ortl 124 Ortl 125 Ortl 126 Ortl 127 Ortl 128 Ortl 129 Ortl 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	rtho rtho rtho rtho rtho rtho nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	Reamer Rongeur	888.4540 888.4540 888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HTO HTX HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 1 1 1 1 2 1	CCCCCC N	Y Y Y Y Y N N
124 Ortl 125 Ortl 126 Ortl 127 Ortl 128 Ortl 129 Ortl 130 Phy 131 Phy 132 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138 Res	rtho rtho rtho rtho rtho edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Rongeur Scissors Staple Driver Trephine Flexible Reamers/Drills External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	888.4540 888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HTX HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 1 1 2 1 1	CCCCC N	Y Y Y Y N N
125 Ortil 126 Ortil 127 Ortil 128 Ortil 129 Ortil 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	rtho rtho rtho rtho edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Scissors	888.4540 888.4540 888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HRR HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 1 2 1 1	C C C C C C C N N	Y Y Y N N
126 Ortil 127 Ortil 128 Ortil 129 Ortil 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138 Res	rtho rtho rtho edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Staple Driver Trephine Flexible Reamers/Drills External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	888.4540 888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HXJ HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 2 1 1	C C C N N	Y Y Y N
127 Ortil 128 Ortil 129 Ortil 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	rtho rtho edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Trephine Flexible Reamers/Drills External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	888.4540 886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HWK GEY, HRG JEC KTW KTT NHN IKD	1 1 2 1 1	C C N	Y Y N
128 Orth 129 Orth 130 Phy 131 Phy 132 Phy 133 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	rtho rtho edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Flexible Reamers/Drills External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	886.4070 878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	GEY, HRG JEC KTW KTT NHN IKD	1 2 1 1	C N N	Y N N
129 Orti 130 Phy Mer 131 Phy Mer 132 Phy Mer 133 Phy Mer 135 Phy Mer 135 Phy Mer 136 Phy Mer 136 Phy Mer 137 Rac 138 Res	rtho edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	 External Fixation Frame Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Mechanical Wrist. 	878.4820 888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11 11 1	HRG JEC KTW KTT NHN IKD	2 1 1	N N	N N
130 Phy Mei 131 Phy Mai 132 Phy Mei 133 Phy Mei 135 Phy Mei 136 Pha Sur 137 Rac 138 Res	nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	888.3040 888.3030 Unclassified 890.1175 890.3420 890.3420	11	JEC KTW KTT NHN IKD	1	N	N
130 Phy Mei 131 Phy Mai 132 Phy Mei 133 Phy Mei 135 Phy Mei 136 Pha Sur 137 Rac 138 Res	nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine nysical edicine	Non-Heating Lamp for Adjunctive Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	888.3030 Unclassified 890.1175 890.3420 890.3420	11	KTW KTT NHN IKD	1	N	N
Mei 131 Phy 132 Phy 132 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138	edicine hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Use Inpatient Therapy. Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	890.1175 890.3420 890.3420	I	NHN IKD	1		
131 Phy 132 Phy 133 Phy 133 Phy 134 Phy 135 Phy 136 Pla: Sur 138 Res 138	hysical edicine hysical edicine hysical edicine hysical edicine hysical edicine	Electrode Cable, External Limb Component, Hip Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	890.3420 890.3420	I			N	Y
132 Phy 133 Phy 134 Phy 135 Phy 135 Phy 136 Pla: Sur 137 Rac 138 Res	nysical edicine nysical edicine nysical edicine nysical edicine	Joint. External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.	890.3420		ISL	2		
133 Phy 134 Phy 135 Phy 135 Phy 136 Pla: Sur 137 Rac 138 Res	nysical edicine nysical edicine nysical edicine	External Limb Component, Knee Joint. External Limb Component, Me- chanical Wrist.					N	Y
134 Phy 135 Phy 136 Pla: Sur 137 138 Res	nysical edicine nysical edicine	External Limb Component, Me- chanical Wrist.		1	ISY	2	N	Y
135 Phy Mer 136 Pla Sur 137 Rac 138 Res	nysical edicine		890.3420	I	ISZ	2	N	Y
136 Pla Sur 137 Rac 138 Res		· · · ·	890.3420	I	IQQ	2	N	Y
137 Rao 138 Res	Irgon/	der Joint. Stapler	878.4800	I	GAG,	2	с	Y
138 Res	urgery adiology	Isotope Needle	892.5730	11	GEF, FHM, HBT IWF	3	С	N
_	0,		Unclassified		LNZ	3	c	N
139 Res	esp	Endotracheal Tube Changer			BSP	3	c	1
140 Doc	esp	Anesthesia conduction needle	868.5150		MIA	3	c	N
	esp esp	Short term spinal needle Respiratory Therapy And Anes- thesia Breathing Circuits.	868.5150 868.5240	 	CAI	2	s	N Y
142 Res	esp	Oral And Nasal Catheters	868.5350	1	BZB	1	с	Y
	esp	Gas Masks	868.5550		BSJ	1	S	Y
			868.5620		BYP	1	N	Y
		Breathing Mouthpiece	868.5730		BTR	3	C	N
	esp	Tracheal Tube	868.5810		BZA	2	s	Y
					BZD	3	S	N
	esp	CPAP Mask	868.5905					
		Emergency Manual Resuscitator	868.5915		BTM	2	S	N
	esp	Tracheobronchial Suction Catheter	868.6810		BSY	3	S	Y
	urgery	AC-powered Orthopedic Instru- ment and accessories.	Unclassified		HWE	2	c	N
-	urgery	Breast Implant Mammary Sizer	Unclassified		MRD	1	C	N
-	urgery	Ultrasonic Surgical Instrument	Unclassified		LFL	3	С	N
153 Sur	urgery	Trocar	874.4420	1	KAB, KBG,	3	С	Y
154 Sur	urgery	Endoscopic Blades	876.1500	П	KCI GCP, GCR	2	с	N
155 Sur	urgery	Endoscopic Guidewires	876.1500	II	GCP, GCR	1	с	N
-	urgery urgery	Inflatable External Extremity Splint Noninflatable External Extremity Splint.	878.3900 878.3910	1	FZF FYH	1 1	N N	Y Y
158 Sur	urgery	Catheter needle	878.4200	1	GCB	3	с	Y
	urgery	Implantable Clip	878.4300	ii ii	FZP	3	c	Ň
-	urgery	Electrosurgical And Coagulation Unit With Accessories.	878.4400		BWA	2	c	N
161 Sur	urgery	Electrosurgical Apparatus	878.4400	п	НАМ	2	с	N
-	urgery	Electrosurgical Cutting & Coagula- tion Device & Accessories.	878.4400	11	GEI	2	c	N
163 Sur	urgery	Electrosurgical Device	878.4400	п	DWG	2	с	N
-		Electrosurgical Electrode	878.4400	ii ii	JOS	2	c	N
-	urgery	Implantable Staple, Clamp, Clip for Suturing Apparatus.	878.4750	II	GDW	3	c	N
166 Sur	urgery	Percutaneous biopsy device	878.4800	1	MJG	3	С	Y
167 Sur	• •	Gastro-Urology needle	878.4800		FHR	3	c	Ý

ATTACHMENT 1.—LIST OF SUDS KNOWN TO BE REPROCESSED OR CONSIDERED FOR REPROCESSING—Continued

	Medical specialty	Device type	Regulation No.	Class	Product code	Risk 1,2,3,3*	Critical/semi- critical/non- critical	Premarket exempt
168	Surgery	Aspiration and injection needle	878.4800	1	GAA	3	С	Y
169	Surgery	Biopsy Brush	878.4800	1	GEE	1	C	Y
170	Surgery	Blood Lancet	878.4800	1	FMK	1	C	Y
171	Surgery	Bone Hook	878.4800	1	KIK	1	С	Y
172	Surgery	Cardiovascular Biopsy Needle	878.4800	1	DWO	3	C	Y
173	Surgery	Clamp	878.4800	1	GDJ	1	C	Y
174	Surgery	Clamp	878.4800	1	HXD	1	C	Y
175	Surgery	Curette	878.4800		HTF	1	C	Y
176	Surgery	Disposable Surgical Instrument	878.4800		KDC	1	C	Y
177 178	Surgery	Disposable Vein Stripper	878.4800 878.4800		GAJ GDI	1	C C	Y Y
178	Surgery	Dissector Forceps	878.4800		GEN	2	c	Y
180	Surgery	Forceps	878.4800	li	HTD	2	č	Ý
181	Surgery	Gouge	878.4800	i	GDH	1	č	Ý
182	Surgery	Hemostatic Clip Applier	878.4800	1	HBT	2	C	Y
183	Surgery	Hook	878.4800	1	GDG	1	C	Y
184	Surgery	Manual Instrument	878.4800	1	MDM,	1	C	Y
	_				MDW		_	
185	Surgery	Manual Retractor	878.4800		GZW	1	C	Y
186	Surgery	Manual Saw And Accessories	878.4800	1	GDR	1	С	Y
187	Surgony	Manual Saw And Accessories	070 4000		HAC HAC	1	с	Y
188	Surgery	Manual Surgical Chisel	878.4800 878.4800		FZO	1	c	Y
189	Surgery	Mastoid Chisel	878.4800		JYD	1	c	Ý
190	Surgery	Orthopedic Cutting Instrument	878.4800		HTZ	1	č	Ý
191	Surgery	Orthopedic Spatula	878.4800	i	HXR	1	c	Ŷ
192	Surgery	Osteotome	878.4800	1	HWM	1	С	Y
193	Surgery	Rasp	878.4800	1	GAC	1	C	Y
194	Surgery	Rasp	878.4800	1	HTR	1	C	Y
195	Surgery	Retractor	878.4800		GAD	1	C	Y
196	Surgery	Retractor	878.4800		HXM		C	Y
197 198	Surgery	Saw	878.4800 878.4800		HSO GES	1	C C	Y Y
198	Surgery	Scalpel Blade	878.4800		GDZ	1	c	Y
200	Surgery	Scissors	878.4800		LRW	1	c	Ý
201	Surgery	Snare	878.4800	li	GAE	1	č	Ý
202	Surgery	Spatula	878.4800	1	GAF	1	С	Y
203	Surgery	Staple Applier	878.4800	1	GEF	2	C	Y
204	Surgery	Stapler	878.4800	1	GAG	2	C	Y
205	Surgery	Stomach And Intestinal Suturing Apparatus.	878.4800		FHM	2	C	Y
206 207	Surgery	Surgical Curette	878.4800 878.4800		FZS FZT	1	C C	Y Y
207	Surgery	Surgical Knife	878.4800		EMF	1	S	Y
209	Surgery	Laser Powered Instrument	878.4810	ii ii	GEX	2	c	Ň
210	Surgery	Ac-Powered Motor	878.4820	1	GEY	2	C	Y
211	Surgery	Bit	878.4820	1	GFG	1	С	Y
212	Surgery	Bur	878.4820	1	GFF,	1	C	Y
213	Surgery	Cardiovascular Surgical Saw Blade.	878.4820	I	GEY DWH	1	с	Y
214	Surgery	Chisel (Osteotome)	878.4820	1	KDG	1	С	Y
215	Surgery	Dermatome	878.4820	1	GFD	1	c	Ý
216	Surgery	Electrically Powered Saw	878.4820	1	DWI	2	С	Y
217	Surgery	Pneumatic Powered Motor	878.4820		GET	2	C	Y
218	Surgery	Pneumatically Powered Saw	878.4820		KFK	2	C	Y
219	Surgery	Powered Saw And Accessories	878.4820		HAB	2	C	Y Y
220 221	Surgery	Saw Blade Nonpneumatic Tourniquet	878.4820 878.5900		GFA GAX	1	C N	Y Y
221	Surgery	Pneumatic Tourniquet	878.5900		KCY	1	N	Y Y
223	Surgery	Enodoscopic Staplers	888.4540	li	HXJ	2	C	Ý
224	Surgery	Trocar	876.1500	ii ii	GCJ,	3	č	Ň
	5,		870.1390		DRC			
225	Surgery	Surgical Cutting Accessories	878.4800,	1	GDZ,	2	С	Y
			874.4420		GDX, GES, KBQ, KAS			
226	Surgery	Electrosurgical Electrodes/Han-	876.4300	п	HAM,	2	с	N
		dles/Pencils.	878.4400		GEI,		-	
					FAS			

ATTACHMENT 1.—LIST OF SUDS KNOWN TO BE R	REPROCESSED OR CONSIDERED	FOR REPROCESSING—Continued
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	Medical specialty	Device type	Regulation No.	Class	Product code	Risk 1,2,3,3*	Critical/semi- critical/non- critical	Premarket exempt
227	Surgery	Scissor Tips	878.4800, 884.4520, 874.4420	1	LRW, HDK, HDJ, JZB, KBD	2	С	Y
228	Surgery	Laser Fiber Delivery Systems	878.4810 874.4500 886.4390 884.4550 886.4690	II.	GEX EWG LLW HQF HHR HQB	1	С	Ν

1 = low risk according to RPS

2 = moderate risk according to RPS 3 = high risk according to RPS

 3^* = high risk due to neurological use

Dated: April 23, 2003.

Jeffrey Shuren,

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of Inspector General

Publication of OIG Special Advisory Bulletin on Contractual Joint Ventures

AGENCY: Office of Inspector General (OIG), HHS.

ACTION: Notice.

SUMMARY: The OIG periodically develops and issues guidance, including Special Advisory Bulletins, to alert and inform the health care industry about potential problems or areas of special interest. This **Federal Register** notice sets forth the recently issued OIG Special Advisory Bulletin addressing certain contractual joint venture arrangements.

FOR FURTHER INFORMATION CONTACT:

Vicki Robinson or Joel Schaer, Office of Counsel to the Inspector General, (202) 619–0335.

SUPPLEMENTARY INFORMATION:

Special Advisory Bulletin: Contractual Joint Ventures (April 2003)

Introduction

This Special Advisory Bulletin addresses certain complex contractual arrangements for the provision of items and services previously identified as suspect in our 1989 Special Fraud Alert on Joint Venture Arrangements.¹ While much of the discussion in the 1989 Special Fraud Alert focused on investor referrals to newly formed entities, we observed that:

[t]he Office of Inspector General has become aware of a proliferation of arrangements between those in a position to refer business, such as physicians, and those providing items or services for which Medicare or Medicaid pays. Some examples of the items or services provided in these arrangements include clinical diagnostic laboratory services, durable medical equipment (DME), and other diagnostic services. Sometimes these deals are called "joint ventures." A joint venture may take a variety of forms: it may be a contractual arrangement between two or more parties to cooperate in providing services, or it may involve the creation of a new legal entity by the parties, such as a limited partnership or closely held corporation, to provide such services. (Emphasis added.)

Notwithstanding that caution, the Office of Inspector General (OIG) is concerned that contractual joint venture arrangements are proliferating.²

A. Questionable Contractual Arrangements

The federal anti-kickback statute, section 1128B(b) of the Social Security Act (the Act), prohibits knowingly and willfully soliciting, receiving, offering, or paying anything of value to induce referrals of items or services payable by a federal health care program. Kickbacks

also available on our Web page at http:// oig.hhs.gov/fraud/docs/alertsandbulletins/ 121994.html.

² The kinds of contractual arrangements addressed in this Special Advisory Bulletin are sometimes referred to as "joint ventures" or "contractual joint ventures" or may be referenced by other terminology. For purposes of the analysis set forth in this Bulletin, a "joint venture" is any common enterprise with mutual economic benefit. The application of this Bulletin is not limited to "joint ventures" that meet technical qualifications under applicable state or common law.

are harmful because they can (1) distort medical decision-making, (2) cause overutilization, (3) increase costs to the federal health care programs, and (4) result in unfair competition by freezing out competitors unwilling to pay kickbacks. Both parties to an impermissible kickback transaction may be liable. Violation of the statute constitutes a felony punishable by a maximum fine of \$25,000, imprisonment up to 5 years, or both. The OIG may also initiate administrative proceedings to exclude persons from the federal health care programs or to impose civil money penalties for kickback violations under sections 1128(b)(7) and 1128A(a)(7) of the Act.

This Special Advisory Bulletin focuses on questionable contractual arrangements where a health care provider in one line of business (hereafter referred to as the "Owner") expands into a related health care business by contracting with an existing provider of a related item or service (hereafter referred to as the "Manager/ Supplier") to provide the new item or service to the Owner's existing patient population, including federal health care program patients. The Manager/ Supplier not only manages the new line of business, but may also supply it with inventory, employees, space, billing, and other services. In other words, the Owner contracts out substantially the entire operation of the related line of business to the Manager/Supplierotherwise a potential competitorreceiving in return the profits of the business as remuneration for its federal program referrals.

Some examples of potentially problematic contractual arrangements include the following:

¹ The 1989 Special Fraud Alert was reprinted in the **Federal Register** in 1994. *See* 59 FR 65372 (December 19, 1994). The Special Fraud Alert is