§ 890.5900 Power traction equipment.

(a) Identification. Powered traction equipment consists of powered devices intended for medical purposes for use in conjunction with traction accessories, such as belts and harnesses, to exert therapeutic pulling forces on the patient’s body.

(b) Classification. Class II (performance standards).

§ 890.5925 Traction accessory.

(a) Identification. A traction accessory is a nonpowered accessory device intended for medical purposes to be used with powered traction equipment to aid in exerting therapeutic pulling forces on the patient’s body. This generic type of device includes the pulley, strap, head halter, and pelvic belt.

(b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter, subject to the limitations in § 890.9.

§ 890.5940 Chilling unit.

(a) Identification. A chilling unit is a refrigerative device intended for medical purposes to chill and maintain cold packs at a reduced temperature.

(b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter, subject to the limitations in § 890.9.

§ 890.5950 Powered heating unit.

(a) Identification. A powered heating unit is a device intended for medical purposes that consists of an encased cabinet containing hot water and that is intended to heat and maintain hot packs at an elevated temperature.

(b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter, subject to the limitations in § 890.9.

§ 890.5975 Therapeutic vibrator.

(a) Identification. A therapeutic vibrator is an electrically powered device intended for medical purposes that incorporates various kinds of pads and that is held in the hand or attached to the hand or to a table. It is intended for various uses, such as relieving minor aches and pains.

(b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter, subject to the limitations in § 890.9.

PART 892—RADIOLOGY DEVICES

Subpart A—General Provisions

Sec.
892.1 Scope.
892.3 Effective dates of requirement for premarket approval.
892.9 Limitations of exemptions from section 510(k) of the Federal Food, Drug, and Cosmetic Act (the act).

Subpart B—Diagnostic Devices

892.1000 Magnetic resonance diagnostic device.
892.1100 Scintillation (gamma) camera.
892.1110 Positron camera.
892.1130 Nuclear whole body counter.
892.1170 Bone densitometer.
892.1180 Bone sonometer.
892.1200 Emission computed tomography system.
892.1220 Fluorescent scanner.
892.1300 Nuclear rectilinear scanner.
892.1310 Nuclear tomography system.
892.1320 Nuclear uptake probe.
892.1330 Nuclear whole body scanner.
892.1350 Nuclear scanning bed.
892.1360 Radionuclide dose calibrator.
892.1370 Nuclear anthropomorphic phantom.
892.1380 Nuclear flood source phantom.
892.1390 Radionuclide rebreathing system.
892.1400 Nuclear sealed calibration source.
892.1410 Nuclear electrocardiograph synchronizer.