(b) **Iontophoresis device intended for any other purposes**—

(1) **Identification.** An iontophoresis device is a device that is intended to use a direct current to introduce ions of soluble salts or other drugs into the body for medical purposes other than those specified in paragraph (a) of this section.

(2) **Classification.** Class III (premarket approval).

(c) **Date PMA or notice of completion of a PDP is required.** No effective date has been established of the requirement for premarket approval for the device described in paragraph (b)(1). See §890.3.


§ 890.5575 **Powered external limb overload warning device.**

(a) **Identification.** A powered external limb overload warning device is a device intended for medical purposes to warn a patient of an overload or an underload in the amount of pressure placed on a leg.

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

§ 890.5650 **Powered inflatable tube massager.**

(a) **Identification.** A powered inflatable tube massager is a powered device intended for medical purposes to relieve minor muscle aches and pains and to increase circulation. It simulates kneading and stroking of tissues with the hands by use of an inflatable pressure cuff.

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

§ 890.5660 **Therapeutic massager.**

(a) **Identification.** A therapeutic massager is an electrically powered device intended for medical purposes, such as to relieve minor muscle 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.


§ 890.5700 **Cold pack.**

(a) **Identification.** A cold pack is a device intended for medical purposes that consists of a compact fabric envelope containing a specially hydrated pliable silicate gel capable of forming to the contour of the body and that provides cold therapy for body surfaces.

(b) **Classification.** Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807. The device also is exempt from the current good manufacturing practice requirements of the quality system regulation in part 820, with the exception of §820.180, with respect to general requirements concerning records, and §820.198, with respect to complaint files.

§ 890.5710 **Hot or cold disposable pack.**

(a) **Identification.** A hot or cold disposable pack is a device intended for medical purposes that consists of a sealed plastic bag incorporating chemicals that, upon activation, provides hot or cold therapy for body surfaces.

(b) **Classification.** Class I (general controls). Except when intended for use on infants, the device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to §890.9.


§ 890.5720 **Water circulating hot or cold pack.**

(a) **Identification.** A water circulating hot or cold pack is a device intended for medical purposes that operates by pumping heated or chilled water through a plastic bag and that provides hot or cold therapy for body surfaces.

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


§ 890.5730 **Moist heat pack.**

(a) **Identification.** A moist heat pack is a device intended for medical purposes that consists of silica gel in a