## Food and Drug Administration, HHS

subpart E of part 807 of this chapter, subject to the limitations in §878.9.

[53 FR 23872, June 24, 1988, as amended at 61 FR 1123, Jan. 16, 1996; 66 FR 38803, July 25, 2001]

## §878.5360 Tweezer-type epilator.

(a) *Identification*. The tweezer-type epilator is an electrical device intended to remove hair. The energy provided at the tip of the tweezer used to remove hair may be radio frequency, galvanic (direct current), or a combination of radio frequency and galvanic energy.

(b) *Classification*. Class I (general controls). The device is exempt from premarket notification procedures in subpart E of part 807 of this chapter subject to §878.9.

[63 FR 57060, Oct. 26, 1998]

# §878.5400 Low level laser system for aesthetic use

(a) *Identification*. A Low Level Laser System for Aesthetic Use is a device using low level laser energy for the disruption of adipocyte cells within the fat layer for the release of fat and lipids from these cells for noninvasive aesthetic use.

(b) Classification. Class II (special controls). The special control for this device is the FDA guidance document entitled "Guidance for Industry and Food and Drug Administration Staff; Class II Special Controls Guidance Document: Low Level Laser System for Aesthetic Use." See §878.1(e) for the availability of this guidance document.

[76 FR 20842, Apr. 14, 2011]

# §878.5650 Topical oxygen chamber for extremities.

(a) *Identification*. A topical oxygen chamber for extremities is a device that is intended to surround a patient's limb and apply humidified oxygen topically at a pressure slightly greater than atmospheric pressure to aid healing of chronic skin ulcers such as bedsores.

(b) *Classification*. Class II (special controls). The special control for this device is FDA's "Class II Special Controls Guidance: Topical Oxygen Chamber for Extremities." See §878.1(e) for

the availability of this guidance document.

[76 FR 22807, Apr. 25, 2011]

### §878.5900 Nonpneumatic tourniquet.

(a) *Identification*. A nonpneumatic tourniquet is a device consisting of a strap or tubing intended to be wrapped around a patient's limb and tightened to reduce circulation.

(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 §878.9.

[53 FR 23872, June 24, 1988, as amended at 54
FR 13828, Apr. 5, 1989; 59 FR 63010, Dec. 7, 1994; 66 FR 38803, July 25, 2001]

#### §878.5910 Pneumatic tourniquet.

(a) *Identification*. A pneumatic tourniquet is an air-powered device consisting of a pressure-regulating unit, connecting tubing, and an inflatable cuff. The cuff is intended to be wrapped around a patient's limb and inflated to reduce or totally occlude circulation during surgery.

(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 §878.9.

[53 FR 23872, June 24, 1988, as amended at 61 FR 1123, Jan. 16, 1996; 66 FR 38803, July 25, 2001]

# PART 880—GENERAL HOSPITAL AND PERSONAL USE DEVICES

### Subpart A—General Provisions

Sec. 880.1 Scope

- 880.3 Effective dates of requirement for premarket approval.
- 880.9 Limitations of exemptions from section 510(k) of the Federal Food, Drug, and Cosmetic Act (the act).

## Subpart B [Reserved]

#### Subpart C—General Hospital and Personal Use Monitoring Devices

- 880.2200 Liquid crystal forehead temperature strip.
- 880.2400 Bed-patient monitor.
- 880.2420 Electronic monitor for gravity flow infusion systems.

### Pt. 880