§ 886.3300 Absorbable implant (scleral buckling method).
(a) Identification. An absorbable implant (scleral buckling method) is a device intended to be implanted on the sclera to aid retinal reattachment.
(b) Classification. Class II.

§ 886.3320 Eye sphere implant.
(a) Identification. An eye sphere implant is a device intended to be implanted in the eyeball to occupy space following the removal of the contents of the eyeball with the sclera left intact.
(b) Classification. Class II.

§ 886.3340 Extraocular orbital implant.
(a) Identification. An extraocular orbital implant is a nonabsorbable device intended to be implanted during scleral surgery for buckling or building up the floor of the eye, usually in conjunction with retinal reattachment. Injectable substances are excluded.
(b) Classification. Class II.

§ 886.3400 Keratoprosthesis.
(a) Identification. A keratoprosthesis is a device intended to provide a transparent optical pathway through an opacified cornea, either intraoperatively or permanently, in an eye that is not a reasonable candidate for a corneal transplant.
(b) Classification. Class II. The special controls for this device are FDA’s:
(2) “510(k) Sterility Review Guidance of 2/12/90 (K90–1),” and
(3) “Guidance on 510(k) Submissions for Keratoprostheses.”
[65 FR 17147, Mar. 31, 2000, as amended at 66 FR 18542, Apr. 10, 2001]

§ 886.3600 Intraocular lens.
(a) Identification. An intraocular lens is a device made of materials such as glass or plastic intended to be implanted to replace the natural lens of an eye.
(b) Classification. Class III.
(c) Date PMA or notice of completion of a PDP is required. As of May 28, 1976, an approval under section 515 of the act is required before this device may be commercially distributed. See §886.3.

§ 886.3800 Scleral shell.
(a) Identification. A scleral shell is a device made of glass or plastic that is intended to be inserted for short time periods over the cornea and proximal cornea sclera for cosmetic or reconstructive purposes. An artificial eye is usually painted on the device. The device is not intended to be implanted.
(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 §886.9.


§ 886.3920 Aqueous shunt.
(a) Identification. An aqueous shunt is an implantable device intended to reduce intraocular pressure in the anterior chamber of the eye in patients with neovascular glaucoma or with glaucoma when medical and conventional surgical treatments have failed.
(b) Classification. Class II. The special controls for this device are FDA’s:
(2) “510(k) Sterility Review Guidance of 2/12/90 (K90–1),” and
(3) “Aqueous Shunts—510(k) Submissions.”

[65 FR 17147, Mar. 31, 2000, as amended at 66 FR 18542, Apr. 10, 2001]

Subpart E—Surgical Devices

§ 886.4070 Powered corneal burr.
(a) Identification. A powered corneal burr is an AC-powered or battery-powered device that is a motor and drilling tool intended to remove rust rings from the cornea of the eye.
(b) Classification. Class I (general controls). When intended only for rust ring removal, the device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to §886.9.