(2) Labeling:
(i) Indication: Only to evaluate the endometrium.
(ii) Contraindications: Pregnancy, history of uterine perforation, or a recent cesarean section, and
(iii) Warning: Do not attach to a wall or any external suction, and
(3) Design and Testing:
(i) The sampling component is covered within the vagina, and
(ii) Intrauterine pressure should not exceed 50 millimeters of mercury.

§ 884.1300 Uterotubal carbon dioxide insufflator and accessories.
(a) Identification. A uterotubal carbon dioxide insufflator and accessories is a device used to test the patency (lack of obstruction) of the fallopian tubes by pressurizing the uterus and fallopian tubes and filling them with carbon dioxide gas.
(b) Classification. Class II (performance standards).

§ 884.1425 Perineometer.
(a) Identification. A perineometer is a device consisting of a fluid-filled sack for intravaginal use that is attached to an external manometer. The devices measure the strength of the perineal muscles by offering resistance to a patient’s voluntary contractions of these muscles and is used to diagnose and to correct, through exercise, urinary incontinence or sexual dysfunction.
(b) Classification. Class II (performance standards).

§ 884.1550 Amniotic fluid sampler (amniocentesis tray).
(a) Identification. The amniotic fluid sampler (amniocentesis tray) is a collection of devices used to aspirate amniotic fluid from the amniotic sac via a transabdominal approach. Components of the amniocentesis tray include a disposable 3 inch 20 gauge needle with stylet and a 30 cc. syringe, as well as the various sample collection accessories, such as vials, specimen containers, medium, drapes, etc. The device is used at 16–18 weeks gestation for antepartum diagnosis of certain congenital abnormalities or anytime after 24 weeks gestation when used to assess fetal maturity.
(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 §884.9.

§ 884.1600 Transabdominal amnioscope (fetoscope) and accessories.
(a) Identification. A transabdominal amnioscope is a device designed to permit direct visual examination of the fetus by a teleoscopic system via abdominal entry. The device is used to ascertain fetal abnormalities, to obtain fetal blood samples, or to obtain fetal tissue. This generic type of device may include the following accessories: trocar and cannula, instruments used through an operating channel or through a separate cannula associated with the amnioscope, light source and cables, and component parts.
(b) Classification. Class III (premarket approval).
(c) Date premarket approval application (PMA) or notice of completion of a product development protocol (PDP) is required. A PMA or a notice of completion of a PDP is required to be filed with the Food and Drug Administration on or before January 29, 1987 for any transabdominal amnioscope (fetoscope) and accessories that was in commercial distribution before May 28, 1976, or that has on or before January 29, 1987 been found to be substantially equivalent to a transabdominal amnioscope (fetoscope) and accessories that was in commercial distribution before May 28, 1976. Any other transabdominal amnioscope (fetoscope) and accessories shall have an approved PMA or a declared completed PDP in