Food and Drug Administration, HHS

subpart E of part 807 of this chapter subject to the limitations in §872.9. If the device is not labeled or otherwise represented as sterile, it is exempt from the current good manufacturing practice requirements of the quality system regulation in part 820 of this chapter, with the exception of §820.180, with respect to general requirements concerning records, and §820.198, with respect to complaint files.

[52 FR 30097, Aug. 12, 1987, as amended at 54 FR 13831, Apr. 5, 1989; 66 FR 38800, July 25, 2001]

§872.6880 Preformed impression tray.

(a) Identification. A preformed impression tray is a metal or plastic device intended to hold impression material, such as alginate, to make an impression of a patient's teeth or alveolar process (bony tooth sockets) to reproduce the structure of a patient's teeth and gums.

(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 §872.9. If the device is not labeled or otherwise represented as sterile, it is exempt from the current good manufacturing practice requirements of the quality system regulation in part 820 of this chapter, with the exception of §820.180, with respect to general requirements concerning records, and §820.198, with respect to complaint files.

[52 FR 30097, Aug. 12, 1987, as amended at 54 FR 13832, Apr. 5, 1989; 66 FR 38800, July 25, 2001]

§872.6890 Intraoral dental wax.

(a) Identification. Intraoral dental wax is a device made of wax intended to construct patterns from which custom made metal dental prostheses, such as crowns and bridges, are cast. In orthodontic dentistry, the device is intended to make a pattern of a patient's bite to make study models of the teeth.

(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 §872.9. If the device is not labeled or otherwise represented as sterile, it is exempt from the current good manufacturing

practice requirements of the quality system regulation in part 820 of this chapter, with the exception of §820.180, with respect to general requirements concerning records, and §820.198, with respect to complaint files.

[52 FR 30097, Aug. 12, 1987, as amended at 59 FR 63009, Dec. 7, 1994; 66 FR 38800, July 25, 2001]

PART 874—EAR, NOSE, AND THROAT DEVICES

Subpart A—General Provisions

Sec.

874.1 Scope.

874.3 Effective dates of requirement for premarket approval.

874.9 Limitations of exemptions from section 510(k) of the Federal Food, Drug, and Cosmetic Act (the act).

Subpart B—Diagnostic Devices

 $874.1050 \quad Audiometer.$

874.1060 Acoustic chamber for audiometric testing.

874.1070 Short increment sensitivity index (SISI) adapter.

874.1080 Audiometer calibration set.

874.1090 Auditory impedance tester.

874.1100 Earphone cushion for audiometric testing.

874.1120 Electronic noise generator for audiometric testing.

874.1325 Electroglottograph.

874.1520 Gustometer

874.1600 Olfactory test device.

874.1800 Air or water caloric stimulator.

874.1820 Surgical nerve stimulator/locator.

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Subpart C [Reserved]

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874.3430 Middle ear mold.

874.3450 Partial ossicular replacement prosthesis.

874.3495 Total ossicular replacement prosthesis.

874.3540 Prosthesis modification instrument for ossicular replacement surgery.

874.3620 Ear, nose, and throat synthetic polymer material.