(c) In addition to complying with the requirements of this part, owners or operators of device establishments that manufacture radiation-emitting electronic products, as defined in §1000.3 of this chapter, shall comply with the reporting requirements of part 1002 of this chapter.

[42 FR 42526, Aug. 23, 1977, as amended at 72 FR 73601, Dec. 28, 2007]

§ 807.85 Exemption from premarket notification.

- (a) A custom device is exempt from premarket notification requirements of this subpart if the device is within the meaning of section 520(b) of the Federal Food, Drug, and Cosmetic Act.
- (1) It is intended for use by a patient named in the order of the physician or dentist (or other specially qualified person); or
- (2) It is intended solely for use by a physician or dentist (or other specially qualified person) and is not generally available to, or generally used by, other physicians or dentists (or other specially qualified persons).
- (b) A distributor who places a device into commercial distribution for the first time under his own name and a repackager who places his own name on a device and does not change any other labeling or otherwise affect the device shall be exempted from the premarket notification requirements of this subpart if:
- (1) The device was in commercial distribution before May 28, 1976; or
- (2) A premarket notification submission was filed by another person.

 $[42\ FR\ 42526,\ Aug.\ 23,\ 1977,\ as\ amended\ at\ 81\ FR\ 70340,\ Oct.\ 12,\ 2016]$

§807.87 Information required in a premarket notification submission.

Each premarket notification submission shall contain the following information:

- (a) The device name, including both the trade or proprietary name and the common or usual name or classification name of the device.
- (b) The establishment registration number, if applicable, of the owner or operator submitting the premarket notification submission.
- (c) The class in which the device has been put under section 513 of the act

- and, if known, its appropriate panel; or, if the owner or operator determines that the device has not been classified under such section, a statement of that determination and the basis for the person's determination that the device is not so classified.
- (d) Action taken by the person required to register to comply with the requirements of the act under section 514 for performance standards.
- (e) Proposed labels, labeling, and advertisements sufficient to describe the device, its intended use, and the directions for its use. Where applicable, photographs or engineering drawings should be supplied.
- (f) A statement indicating the device is similar to and/or different from other products of comparable type in commercial distribution, accompanied by data to support the statement. This information may include an identification of similar products, materials, design considerations, energy expected to be used or delivered by the device, and a description of the operational principles of the device.
- (g) Where a person required to register intends to introduce into commercial distribution a device that has undergone a significant change or modification that could significantly affect the safety or effectiveness of the device, or the device is to be marketed for a new or different indication for use, the premarket notification submission must include appropriate supporting data to show that the manufacturer has considered what consequences and effects the change or modification or new use might have on the safety and effectiveness of the device.
- (h) A 510(k) summary as described in §807.92 or a 510(k) statement as described in §807.93.
- (i) A financial certification or disclosure statement or both, as required by part 54 of this chapter.
- (j) For a submission supported by clinical data:
- (1) If the data are from clinical investigations conducted in the United States, a statement that each investigation was conducted in compliance with applicable requirements in the

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protection of human subjects regulations in part 50 of this chapter, the institutional review boards regulations in part 56 of this chapter, or was not subject to the regulations under §56.104 or §56.105, and the investigational device exemptions regulations in part 812 of this chapter, or if the investigation was not conducted in compliance with those regulations, a brief statement of the reason for the noncompliance.

- (2) If the data are from clinical investigations conducted outside the United States, the requirements under §812.28 of this chapter apply. If any such investigation was not conducted in accordance with good clinical practice (GCP) as described in §812.28(a) of this chapter, include either a waiver request in accordance with §812.28(c) of the chapter or a brief statement of the reason for not conducting the investigation in accordance with GCP and a description of steps taken to ensure that the data and results are credible and accurate and that the rights, safety, and wellbeing of subjects have been adequately protected.
- (k) For submissions claiming substantial equivalence to a device which has been classified into class III under section 513(b) of the act:
- (1) Which was introduced or delivered for introduction into interstate commerce for commercial distribution before December 1, 1990; and
- (2) For which no final regulation requiring premarket approval has been issued under section 515(b) of the act, a summary of the types of safety and effectiveness problems associated with the type of devices being compared and a citation to the information upon which the summary is based (class III summary). The 510(k) submitter shall also certify that a reasonable search of all information known or otherwise available about the class III device and other similar legally marketed devices has been conducted (class III certification), as described in §807.94. This information does not refer to information that already has been submitted to the Food and Drug Administration (FDA) under section 519 of the act. FDA may require the submission of the adverse safety and effectiveness data described in the class III summary or citation.

- (1) A statement that the submitter believes, to the best of his or her knowledge, that all data and information submitted in the premarket notification are truthful and accurate and that no material fact has been omitted.
- (m) Any additional information regarding the device requested by the Commissioner that is necessary for the Commissioner to make a finding as to whether or not the device is substantially equivalent to a device in commercial distribution. A request for additional information will advise the owner or operator that there is insufficient information contained in the original premarket notification submission for the Commissioner to make this determination and that the owner or operator may either submit the requested data or a new premarket notification containing the requested information at least 90 days before the owner or operator intends to market the device, or submit a premarket approval application in accordance with section 515 of the act. If the additional information is not submitted within 30 days following the date of the request, the Commissioner will consider the premarket notification to be withdrawn.

(Information collection requirements in this section were approved by the Office of Management and Budget (OMB) and assigned OMB control number 0910–0281)

[42 FR 42526, Aug. 23, 1977, as amended at 57 FR 18066, Apr. 28, 1992; 59 FR 64295, Dec. 14, 1994; 63 FR 5253, Feb. 2, 1998; 83 FR 7385, Feb. 21, 2018]

§ 807.90 Format of a premarket notification submission.

Each premarket notification submission pursuant to this part shall be submitted in accordance with this section. Each submission shall:

- (a)(1) For devices regulated by the Center for Devices and Radiological Health, be addressed to the current address displayed on the website https://www.fda.gov/cdrhsubmissionaddress.
- (2) For devices regulated by the Center for Biologics Evaluation and Research, be addressed to the current address displayed on the website https://www.fda.gov/AboutFDA/CentersOffices/OfficeofMedicalProductsandTobacco/