(1) A use of a food contact substance that is the subject of a regulation in parts 173 through 189 of this chapter; or
(2) A use of a food contact substance that is the subject of an exemption under the threshold of regulation process described in §170.39.

(c) A petition must be submitted under §171.1 of this chapter to authorize the safe use of a food contact substance in either of the following circumstances, unless FDA agrees to accept an FCN for the proposed use.

(1) The use of the food contact substance increases the cumulative dietary concentration to a certain level. For a substance that is a biocide (e.g., it is intended to exert microbial toxicity), this level is equal to or greater than 200 parts per billion in the daily diet (0.6 milligram (mg)/person/day). For a substance that is not a biocide, this level is equal to or greater than 1 part per million in the daily diet (3 mg/person/day); or

(2) There exists a bioassay on the food contact substance, FDA has not reviewed the bioassay, and the bioassay is not clearly negative for carcinogenic effects.

(d) A manufacturer or supplier for which a notification is effective must keep a current address on file with FDA.

(1) The current address may be either the manufacturer’s (or supplier’s) address or the address of the manufacturer’s (or supplier’s) agent.

(2) FDA will deliver correspondence to the manufacturer’s or supplier’s current address.

§ 170.101 Information in a premarket notification for a food contact substance (FCN).

An FCN must contain the following:

(a) A comprehensive discussion of the basis for the manufacturer’s or supplier’s determination that the use of the food contact substance is safe. This discussion must:

(1) Discuss all information and data submitted in the notification; and

(2) Address any information and data that may appear to be inconsistent with the determination that the proposed use of the food contact substance is safe.

(b) All data and other information that form the basis of the determination that the food contact substance is safe under the intended conditions of use. Data must include primary biological data and chemical data.

(c) A good laboratory practice statement for each nonclinical laboratory study, as defined under §58.3(d) of this chapter, that is submitted as part of the FCN, in the form of either:

(1) A signed statement that the study was conducted in compliance with the good laboratory practice regulations under part 58 of this chapter; or

(2) A brief signed statement listing the reason(s) that the study was not conducted in compliance with part 58 of this chapter.

(3) Data from any study conducted after 1978 but not conducted in compliance with part 58 of this chapter must be validated by an independent third party prior to submission to the Food and Drug Administration (FDA), and the report and signed certification of the validating party must be submitted as part of the notification.

(d) Information to address FDA’s responsibility under the National Environmental Policy Act, in the form of either:

(1) A claim of categorical exclusion under §25.30 or §25.32 of this chapter; or

(2) An environmental assessment complying with §25.40 of this chapter.

(e) A completed and signed FDA Form No. 3480.

§ 170.102 Confidentiality of information in a premarket notification for a food contact substance (FCN).

(a) During the 120-day period of the Food and Drug Administration (FDA) review of an FCN, FDA will not disclose publicly any information in that FCN.

(b) FDA will not disclose publicly the information in an FCN that is withdrawn prior to the completion of FDA’s review.

(c) Once FDA completes its review of an FCN, the agency will make its conclusion about the FCN publicly available. For example, if FDA objects to a notification 90 days after the date of receipt, the agency would make available its objection at that time.
(d) By submitting an FCN to FDA, the manufacturer or supplier waives any claim to confidentiality of the information required to adequately describe the food contact substance and the intended conditions of use that are the subject of that FCN.

(e) The following data and information in an FCN are available for public disclosure, unless extraordinary circumstances are shown, on the 121st day after receipt of the notification by FDA, except that no data or information are available for public disclosure if the FCN is withdrawn under §170.103.

1. All safety and functionality data and information submitted with or incorporated by reference into the notification. Safety and functionality data include all studies and tests of a food contact substance on animals and humans and all studies and tests on a food contact substance for establishing identity, stability, purity, potency, performance, and usefulness.

2. A protocol for a test or study, unless it is exempt from disclosure under §20.61 of this chapter.

3. A list of all ingredients contained in a food contact substance, excluding information that is exempt from disclosure under §20.61 of this chapter. Where applicable, an ingredient list will be identified as incomplete.

4. An assay method or other analytical method, unless it serves no regulatory or compliance purpose and is exempt from disclosure under §20.61 of this chapter.

5. All correspondence and written summaries of oral discussions relating to the notification, except information that is exempt for disclosure under §20.61 of this chapter.

6. All other information not subject to an exemption from disclosure under subpart D of part 20 of this chapter.

§ 170.103 Withdrawal without prejudice of a premarket notification for a food contact substance (FCN).

A manufacturer or supplier may withdraw an FCN without prejudice to a future submission to the Food and Drug Administration (FDA) if FDA has not completed review of the FCN. For the purpose of this section, FDA’s review is completed when FDA has allowed 120 days to pass without objecting to the FCN or FDA has issued an objection letter.

§ 170.104 Action on a premarket notification for a food contact substance (FCN).

(a) If the Food and Drug Administration (FDA) does not object to an FCN within the 120-day period for FDA review, the FCN becomes effective.

(b) If an FCN is complete when received, the 120-day review period begins on the date FDA receives the FCN.

1. If any element required under §170.101 is missing from an FCN, then FDA will not accept that FCN and FDA will send an FCN nonacceptance letter to the manufacturer or supplier. If the manufacturer or supplier submits the missing information before FDA sends an FCN nonacceptance letter, the 120-day review period begins on the date of receipt of the missing information.

2. If FDA accepts an FCN, then FDA will acknowledge in writing its receipt of that FCN.

(c) Objection to an FCN:

1. If FDA objects to an FCN, then FDA will send an FCN objection letter. The date of the letter will be the date of FDA’s objection for purposes of section 409(h)(2)(A) of the act.

2. If FDA objects to an FCN within the 120-day period for FDA review, the FCN will not become effective.

3. FDA may object to an FCN if any part of FDA’s 120-day review occurs during a period when this program is not funded as required in section 409(h)(5) of the act.

(d) If FDA and a manufacturer or supplier agree that the notifier may submit a food additive petition proposing the approval of the food contact substance for the use in the manufacturer’s or supplier’s FCN, FDA will consider that FCN to be withdrawn by the manufacturer or supplier on the date the petition is received by FDA.

§ 170.105 The Food and Drug Administration’s (FDA’s) determination that a premarket notification for a food contact substance (FCN) is no longer effective.

(a) If data or other information available to FDA, including data not submitted by the manufacturer or supplier, demonstrate that the intended use of the food contact substance is no