

§516.155

deny the request and notify the requester of FDA's decision in writing.

(b) If FDA denies the request for addition of a new animal drug to the index, FDA shall provide due notice and an opportunity for an informal conference as described in §516.123. A decision of FDA to deny a request to index a new animal drug following an informal conference shall constitute final agency action subject to judicial review.

§516.155 Labeling of indexed drugs.

(a) The labeling of an indexed drug that is found to be eligible for indexing under §516.129(c)(7)(i) shall state, prominently and conspicuously: "*NOT APPROVED BY FDA.—Legally marketed as an FDA indexed product. Extra-label use is prohibited.*" "*This product is not to be used in animals intended for use as food for humans or other animals.*"

(b) The labeling of an indexed drug that was found to be eligible for indexing for use in an early, non-food life stage of a food-producing minor species animal, under §516.129(c)(7)(ii), shall state, prominently and conspicuously: "*NOT APPROVED BY FDA.—Legally marketed as an FDA indexed product. Extra-label use is prohibited.*"

(c) The labeling of an indexed drug shall contain such other information as may be prescribed in the index listing.

§516.157 Publication of the index and content of an index listing.

(a) FDA will make the list of indexed drugs available through the FDA Web site. A printed copy can be obtained by writing to the FDA Freedom of Information Staff or by visiting the FDA Freedom of Information Public Reading Room.

(b) The list will contain the following information for each indexed drug:

- (1) The name and address of the person who holds the index listing;
- (2) The name of the drug and the intended use and conditions of use for which it is indexed;
- (3) Product labeling; and
- (4) Conditions and any limitations that FDA deems necessary regarding use of the drug.

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§516.161 Modifications to indexed drugs.

(a) After a drug is listed in the index, certain modifications to the index listing may be requested. Any modification of an index listing may not cause an indexed drug to be a different drug (or different combination of drugs) or a different dosage form. If such modification is requested, FDA will notify the holder that a new index listing is required for the new drug or dosage form.

(b) Modifications to the indexed drug will fall under one of three categories and must be submitted as follows:

(1) *Urgent changes.* (i) The following modifications to an indexed drug or its labeling should be made as soon as possible, and a request to modify the indexed drug should be concurrently submitted:

(A) The addition to package labeling, promotional labeling, or prescription drug advertising of additional warning, contraindication, side effect, or cautionary information.

(B) The deletion from package labeling, promotional labeling, and drug advertising of false, misleading, or unsupported indications for use or claims for effectiveness.

(C) Changes in manufacturing methods or controls required to correct product or manufacturing defects that may result in serious adverse drug events.

(ii) The modifications described in paragraph (b)(1)(i) of this section must be submitted to the Director, OMUMS, in the form of a request for modification of an indexed drug, and must contain sufficient information to permit FDA to determine the need for the modification and whether the modification appropriately addresses the need.

(iii) FDA will take no action against an indexed drug or index holder solely because modifications of the kinds described in paragraph (b)(1)(i) of this section are placed into effect by the holder prior to receipt of a written notice granting the request if all the following conditions are met:

(A) A request to modify the indexed drug providing a full explanation of the basis for the modifications has been submitted, plainly marked on the mailing cover and on the request as follows: