Food and Drug Administration, HHS

Subpart B—Written Recommendations for Investigations of Orphan Drugs

- 316.10 Content and format of a request for written recommendations.
- 316.12 Providing written recommendations.316.14 Refusal to provide written recommendations.

Subpart C—Designation of an Orphan Drug

- 316.20 Content and format of a request for orphan-drug designation.
- 316.21 Verification of orphan-drug status.
- 316.22 Permanent-resident agent for foreign sponsor.
- 316.23 Timing of requests for orphan-drug designation; designation of already approved drugs.
- 316.24 Deficiency letters and granting orphan-drug designation.
- 316.25 Refusal to grant orphan-drug designation.
- 316.26 Amendment to orphan-drug designation.
- 316.27 Change in ownership of orphan-drug designation.
- 316.28 Publication of orphan-drug designations.
- 316.29 Revocation of orphan-drug designation.
- 316.30 Annual reports of holder of orphandrug designation.

Subpart D—Orphan-Drug Exclusive Approval

- 316.31 Scope of orphan-drug exclusive approval.
- 316.34 FDA recognition of exclusive approval.
- 316.36 Insufficient quantities of orphan drugs.

Subpart E—Open Protocols for Investigations

316.40 Treatment use of a designated orphan drug.

Subpart F—Availability of Information

- 316.50 Guidance documents.
- 316.52 Availability for public disclosure of data and information in requests and applications.

AUTHORITY: 21 U.S.C. 360aa, 360bb, 360cc, 360dd, 371.

Source: 57 FR 62085, Dec. 29, 1992, unless otherwise noted.

EDITORIAL NOTE: Nomenclature changes to part 316 appear at 69 FR 13717, Mar. 24, 2004.

Subpart A—General Provisions

§316.1 Scope of this part.

- (a) This part implements sections 525, 526, 527, and 528 of the act and provides procedures to encourage and facilitate the development of drugs for rare diseases or conditions, including biological products and antibiotics. This part sets forth the procedures and requirements for:
 - (1) Submissions to FDA of:
- (i) Requests for recommendations for investigations of drugs for rare diseases or conditions:
- (ii) Requests for designation of a drug for a rare disease or condition; and
- (iii) Requests for gaining exclusive approval for a drug for a rare disease or condition.
- (2) Allowing a sponsor to provide an investigational drug under a treatment protocol to patients who need the drug for treatment of a rare disease or condition.
- (b) This part does not apply to food, medical devices, or drugs for veterinary use.
- (c) References in this part to regulatory sections of the Code of Federal Regulations are to chapter I of title 21, unless otherwise noted.

[57 FR 62085, Dec. 29, 1992, as amended at 78 FR 35132, June 12, 2013]

§316.2 Purpose.

The purpose of this part is to establish standards and procedures for determining eligibility for the benefits provided for in section 2 of the Orphan Drug Act, including written recommendations for investigations of orphan drugs, a 7-year period of exclusive marketing, and treatment use of investigational orphan drugs. This part is also intended to satisfy Congress' requirements that FDA promulgate procedures for the implementation of sections 525(a) and 526(a) of the act.

§316.3 Definitions.

- (a) The definitions and interpretations contained in section 201 of the act apply to those terms when used in this part.
- (b) The following definitions of terms apply to this part: