

§ 607.20

21 CFR Ch. I (4–1–01 Edition)

Subpart B—Procedures for Domestic Blood Product Establishments

§ 607.20 Who must register and submit a blood product list.

(a) Owners or operators of all establishments, not exempt under section 510(g) of the act or subpart D of this part 607, that engage in the manufacture of blood products are required to register and to submit a list of every blood product in commercial distribution (except that listing information may be submitted by the parent, subsidiary, and/or affiliate company for all establishments when operations are conducted at more than one establishment and there exists joint ownership and control among all the establishments), whether or not the output of such blood product establishment or any particular blood product so listed enters interstate commerce.

(b) Preparatory to engaging in the manufacture of blood products, owners or operators of establishments who are submitting a biologics license application to manufacture blood products are required to register before the biologics license application is approved.

(c) No registration fee is required. Establishment registration and blood product listing do not constitute an admission or agreement or determination that a blood product is a “drug” within the meaning of section 201(g) of the act.

[40 FR 52788, Nov. 12, 1975, as amended at 64 FR 56452, Oct. 20, 1999]

§ 607.21 Times for establishment registration and blood product listing.

The owner or operator of an establishment entering into an operation defined in § 607.3(d) shall register such establishment within 5 days after the beginning of such operation and submit a list of every blood product in commercial distribution at the time. If the owner or operator of the establishment has not previously entered into such operation (defined in § 607.3(d) of this chapter) for which a license is required, registration shall follow within 5 days after the submission of a biologics license application in order to manufacture blood products. Owners or opera-

tors of all establishments so engaged shall register annually between November 15 and December 31 and shall update their blood product listing information every June and December.

[40 FR 52788, Nov. 12, 1975, as amended at 64 FR 56453, Oct. 20, 1999]

§ 607.22 How and where to register establishments and list blood products.

(a) The first registration of an establishment shall be on Form FD-2830 (Blood Establishment Registration and Product Listing) obtainable on request from the Department of Health and Human Services, Food and Drug Administration, Center for Biologics Evaluation and Research (HFB-240), 8800 Rockville Pike, Bethesda, MD 20892, or from Food and Drug Administration district offices. Subsequent annual registration shall also be accomplished on Form FD-2830 which will be furnished by the Food and Drug Administration before November 15 of each year to establishments whose product registration for that year was validated pursuant to § 607.35. The completed form shall be mailed to the above address before December 31 of that year.

(b) The first list of blood products and subsequent June and December updates shall be on Form FD-2830, obtainable upon request as described in paragraph (a) of this section. In lieu of Form FD-2830, tapes for computer input may be submitted if equivalent in all elements of information as specified in Form FD-2830. All formats proposed for such use will require initial review and approval by the Office of Compliance, Center for Biologics Evaluation and Research, Food and Drug Administration.

[40 FR 52788, Nov. 12, 1975, as amended at 49 FR 23833, June 8, 1984; 55 FR 11014, Mar. 26, 1990]

§ 607.25 Information required for establishment registration and blood product listing.

(a) Form FD-2830 (Blood Establishment Registration and Product Listing) requires furnishing or confirming registration information required by the act. This information includes the