product within the prescribed dating period.
(c) Labeling. In addition to the items required by other applicable labeling provisions of this subchapter, the following shall also be included:
(1) Indication of the source of the product immediately following the proper name on both the final container and package label, e.g., human, guinea pig.
(2) Name of the test method(s) recommended for the product on the package label and on the final container label when capable of bearing a full label (see §610.60(a) of this chapter).
(3) A warning on the package label and on the final container label if capable of bearing a full label (see §610.60(a) of this chapter) indicating that the product and antigen if supplied, shall be handled as if capable of transmitting hepatitis.
(4) If the product is dried, the final container label shall indicate "Reconstitution date: " and a statement indicating the period within which the product may be used after reconstitution.
(5) The package shall include a package enclosure providing (i) adequate instructions for use, (ii) a description of all recommended test methods, and (iii) warnings as to possible hazards, including hepatitis, in handling the product and any ancillary reagents and materials accompanying the product.
(d) Final container. A final container shall be sufficiently transparent to permit visual inspection of the contents for presence of particulate matter and increased turbidity. The effectiveness of the contents of a final container shall be maintained throughout its dating period.
(e) Date of manufacture. The date of manufacture of Antibody to Hepatitis B surface Antigen that has been iodinated with radioactive iodine (125I) shall be the day of labeling the antibody with the radionuclide.
(f) Retention samples. Each manufacturer shall retain representative samples of the product in accordance with §600.13 of this chapter except for that which has been iodinated with radioactive iodine. Retention samples of Antibody to Hepatitis B Surface Antigen iodinated with 125I shall consist of a minimum of two complete finished packages of each lot of the diagnostic test kit and shall be retained for a period of at least 90 days from the date of manufacture.
§ 660.3 Reference panel.
A Reference Hepatitis B Surface Antigen Panel shall be obtained from the Center for Biologics Evaluation and Research (HFM–407) (see mailing addresses in §600.2 of this chapter) and shall be used for determining the potency and specificity of Antibody to Hepatitis B Surface Antigen.
§ 660.4 Potency test.
To be satisfactory for release, each filling of Antibody to Hepatitis B Surface Antigen shall be tested against the Reference Hepatitis B Surface Antigen Panel and shall be sufficiently potent to detect the antigen in the appropriate sera of the reference panel by all test methods recommended by the manufacturer in the package insert.
[40 FR 29711, July 15, 1975]
§ 660.5 Specificity.
Each filling of the product shall be specific for antibody to hepatitis B surface antigen, as determined by specificity tests found acceptable by the Director, Center for Biologics Evaluation and Research.
[40 FR 29712, July 15, 1975, as amended at 49 FR 23834, June 8, 1984; 55 FR 11013, Mar. 26, 1990]
§ 660.6 Samples; protocols; official release.
(a) Samples. (1) For the purposes of this section, a sample of product not iodinated with 125I shall be the day of labeling the antibody with the radionuclide.
(b) Retention samples. Each manufacturer shall retain representative samples of the product in accordance with §600.13 of this chapter except for that which has been iodinated with radioactive iodine. Retention samples of Antibody to Hepatitis B Surface Antigen iodinated with 125I shall consist of