2. Administrative and National Policy Requirements:

45 CFR part 74 or 92.

3. Reporting Requirements

Programmatic Reports: Semi-annually and a final report is due 90 days after the end of the grant period.

Financial Reports: Semi-annually and a final report due 90 days after the end of the grant period.

All grantees are required to submit semi-annual program reports; grantees are also required to submit semi-annual financial status reports using the required financial standard form (SF–269). A format for the program report will be sent to all grantees after the awards are made.

VII. Agency Contacts

Program Office Contact: William D. Riley, Family Violence Division, 330 C Street, Rm. 2117, Switzer Building, Washington, DC 20447, E-mail: wriley@acf.hhs.gov, Telephone: (202) 401–5529.

Grants Management Office Contact:
William Wilson, Grants Officer,
Administration on Children, Youth
and Families, Room 2070 Switzer
Building, 330 C Street, SW.,
Washington, DC 20447, (202) 205–
8913, E-mail: wwilson@acf.hhs.gov.

VIII. Other Information

Additional information about this program and its purpose can be located on the following Web site: http://www.acf.hhs.gov/programs/fysb.

Dated: June 9, 2004.

Frank Fuentes,

Deputy Commissioner, Administration on Children, Youth and Families.

[FR Doc. 04–13736 Filed 6–17–04; 8:45 am] BILLING CODE 4184–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Notice of Approval of Abbreviated New Animal Drug Application; Dexamethasone Sodium Phosphate Injection

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration's Center for Veterinary Medicine (CVM) is providing notice that it has approved an original abbreviated new animal drug application (ANADA) filed by Cross Vetpharm Group Ltd. The ANADA provides for the veterinary

prescription use of dexamethasone sodium phosphate injectable solution as a rapid adrenal glucocorticoid and/or anti-inflammatory agent in horses. The applicable sections of the regulations did not require amendment.

FOR FURTHER INFORMATION CONTACT:

Lonnie W. Luther, Center for Veterinary Medicine (HFV–104), Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855, 301–827–8549, email: *lluther@cvm.fda.gov*.

SUPPLEMENTARY INFORMATION: In accordance with section 512(i) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360b(i)) and 21 CFR 514.105(a) and 514.106(a), CVM is providing notice that it has approved original ANADA 200–317 filed by Cross Vetpharm Group Ltd., Broomhill Rd., Tallaght, Dublin 24, Ireland. ANADA 200-317 provides for the veterinary prescription use of DEXIUM-SP (dexamethasone sodium phosphate) Injection as a rapid adrenal glucocorticoid and/or anti-inflammatory agent in horses. Cross Vetpharm Group's DEXIUM-SP Injection is approved as a generic copy of Steris Laboratories, Inc.'s Dexamethasone Injection, approved under NADA 104-606. The ANADA is approved as of April 29, 2004. The basis of approval is discussed in the freedom of information summary.

In accordance with the freedom of information provisions of 21 CFR part 20 and 21 CFR 514.11(e)(2)(ii), a summary of safety and effectiveness data and information submitted to support approval of this application may be seen in the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, between 9 a.m. and 4 p.m., Monday through Friday.

The agency has determined under 21 CFR 25.33(a)(1) that this action is of a type that does not individually or cumulatively have a significant effect on the human environment. Therefore, neither an environmental assessment nor an environmental impact statement is required.

Dated: June 4, 2004.

Stephen F. Sundlof,

Director, Center for Veterinary Medicine. [FR Doc. 04–13790 Filed 6–17–04; 8:45 am] BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Blood Products Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration,

HHS.

ACTION: Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Blood Products Advisory Committee.

General Function of the Committee: To provide advice and recommendations to the agency on FDA's regulatory issues.

Date and Time: The meeting will be held on July 22, 2004, from 8 a.m. to 6 p.m. and on July 23, 2004, from 8 a.m. to 3 p.m.

Location: Holiday Inn Gaithersburg, Two Montgomery Village Ave., Gaithersburg, MD 20877.

Contact Person: Linda A. Smallwood, Center for Biologics Evaluation and Research (HFM–302), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852, 301–827–3514, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), code 3014519516. Please call the Information Line for up-to-date information on this meeting.

Agenda: On July 22, 2004, the committee will hear updates on: FDA current thinking on transfusion related acute lung inflammation (TRALI), and donor blood pressure determination. The committee will also discuss and provide recommendations on the dating of irradiated blood. In the afternoon, the committee will discuss and provide recommendations on the new standards for platelet evaluation and experience with monitoring of bacterial contamination of platelets. On July 23, 2004, the committee will hear an update on West Nile Virus. The committee will also hear presentations, discuss and provide recommendations on hepatitis B virus nucleic acid testing (HBV NAT) for mini-pools. In the afternoon, there will be an informational presentation on current trends in plasma product manufacturing.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person by July 12, 2004. Oral