

Pursuant to Public Law 92-463, notice is hereby given of a meeting of the Secretary's Advisory Committee on Genetic Testing (SACGT), U.S. Public Health Service. The meeting will be held at the International Trade Center, Polaris Ballroom, 1300 Pennsylvania Avenue, NW, Washington, DC 20004, starting on February 24, 2000, at approximately 9:00 a.m. and will recess at approximately 5:30 p.m. The meeting will reconvene on February 25, 2000, at approximately 8:00 a.m. and will adjourn at approximately 5:00 p.m. The meeting will be open to the public. Attendance by the public will be limited by the space available. The committee will review public comments received in response to A Public Consultation on Oversight of Genetic Tests published in the **Federal Register** on December 1, 1999 (64 FR 67273), and work toward the development of final recommendations on the oversight of genetic testing. A limited period of time will be provided for public comment, and individuals interested in participating in the public comment period should contact Ms. Sarah Carr, SACGT Executive Secretary, as shown below.

Under authority of 42 U.S.C. 217a, Section 222 of the Public Health Service Act, as amended, the Department of Health and Human Services (DHHS) established the SACGT to advise and make recommendations to the Secretary through the Assistant Secretary for Health on all aspects of the development and use of genetic tests. The SACGT is directed to: (1) Recommend policies and procedures for the safe and effective incorporation of genetic technologies into health care; (2) assess the effectiveness of existing and future measures for oversight of genetic tests; (3) and identify research needs related to the Committee's purview.

Further information about the SACGT is available at the following web site: <http://www4.od.nih.gov/oba/sacgt.htm>. If you wish to attend, please register through the web site. A draft meeting agenda will be posted to the web site prior to the meeting. Individuals who wish to provide public comments should notify Ms. Carr, by telephone at 301-496-9838 or E-mail at sc112c@nih.gov as soon as possible and provide a copy of their remarks to Ms. Carr by February 15, 2000. Those who plan to attend the meeting and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify Ms. Carr at 301-496-9838. The SACGT office is located at 6000 Executive Boulevard, Suite 302, Bethesda, Maryland 20892.

Dated: February 3, 2000.

Sarah Carr,

Executive Secretary, SACGT.

[FR Doc. 00-2906 Filed 2-7-00; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Notice of Meeting

Office of the Director, Centers for Disease Control and Prevention (CDC), announces the following meeting:

Name: Guide to Community Preventive Services (GCPS) Task Force Meeting.

Times and Dates: 9 a.m.-5:30 p.m., February 9, 2000; 8 a.m.-3:30 p.m., February 10, 2000.

Place: The Renaissance Waverly Hotel, 2450 Galleria Parkway, Atlanta, Georgia 30339, telephone (770) 953-4500.

Status: Open to the public, limited only by the space available. The meeting room accommodates approximately 40 people.

Purpose: The mission of the Task Force is to develop and publish a Guide to Community Preventive Services, which is based on the best available scientific evidence and current expertise regarding essential public health services and what works in the delivery of those services.

Matters to be Discussed: Agenda items include: a briefing of administrative activities in the Community Guide Branch, recommendation approvals for the Oral Health and Tobacco Chapters, updates for the following chapters: Diabetes, Cancer, Motor Vehicle Occupant Injuries, Mental Health, Physical Activity, Nutrition, Sexual Behavioral, Alcohol, Violence Prevention and Sociocultural Environment, an update on the Economic Evaluation and a discussion of actions items for the next quarter.

Agenda items are subject to change as priorities dictate.

Contact Person for Additional Information: Stephanie Zaza, M.D., M.P.H., Chief, CPS Guide Development Activity, Division of Prevention Research and Analytic Methods, Epidemiology Program Office, CDC, 4770 Buford Highway, M/S K-73, Atlanta, Georgia 30341, telephone 770/488-8189.

Persons interested in reserving a space for this meeting should call 770/488-8189 by close of business on February 7, 2000.

The Director, Management Analysis and Services office has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Carolyn J. Russell,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 00-2899 Filed 2-7-00; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 99N-0671P]

Bestblood, Ltd.; Revocation of U.S. License No. 1116

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the revocation of the establishment license (U.S. License No. 1116) and product licenses (the licenses) issued to Bestblood, Ltd., doing business as Optimum Healthcare, Inc., for the manufacture of Whole Blood, Red Blood Cells, Red Blood Cells Frozen, Whole Blood CPD, Red Blood Cells Deglycerolized, and Whole Blood CPDA-1. Bestblood, Ltd., did not respond to a notice of opportunity for a hearing on a proposal to revoke its licenses.

DATES: The revocation of the establishment license (U.S. License No. 1116) and product licenses is effective February 8, 2000.

FOR FURTHER INFORMATION CONTACT: Joseph L. Okrasinski, Jr., Center for Biologics Evaluation and Research (HFM-17), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD. 20852-1448, 301-827-6210.

SUPPLEMENTARY INFORMATION: FDA is revoking the establishment license (U.S. License No. 1116) and product licenses issued to Bestblood, Ltd., doing business as Optimum Healthcare, Inc., 239 Randall St., San Francisco, CA 94131, for the manufacture of Whole Blood, Red Blood Cells, Red Blood Cells Frozen, Whole Blood CPD, Red Blood Cells Deglycerolized, and Whole Blood CPDA-1. Proceedings to revoke the licenses were initiated because an attempted inspection of the facility by FDA, as required under 21 CFR 600.21, revealed that the firm was no longer in operation.

In a certified, return-receipt letter dated June 16, 1997, FDA notified the firm that the attempt to conduct an inspection at Bestblood, Ltd., 239 Randall St., San Francisco, CA 94131 was unsuccessful because the facility was apparently no longer in operation and requested that the firm notify FDA in writing of the firm's status. This letter was sent to 239 Randall St., San Francisco, CA 94131, and also to P.O. Box 843, Cupertino, CA 95054-0843,