draft guidance document represents the agency's current thinking on submitting Type V Drug Master Files to CBER. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirement of the applicable statutes and regulations.

II. Comments

The draft document is being distributed for comment purposes only and is not intended for implementation at this time. Interested persons may submit to the Dockets Management Branch (address above) written comments regarding this draft guidance document. Submit written or electronic comments to ensure adequate consideration in preparation of the final document by November 21, 2001. Two copies of any comments are to be submitted, except individuals may submit one copy. Comments should be identified with the docket number found in the brackets in the heading of this document. A copy of the document and received comments are available for public examination in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

III. Electronic Access

Persons with access to the Internet may obtain the document at http://www.fda.gov/cber/guidelines.htm.

Dated: August 13, 2001.

Margaret M. Dotzel,

Associate Commissioner for Policy.
[FR Doc. 01–21246 Filed 8–22–01; 8:45 am]
BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. 00D-1618]

"Guidance for Industry: Variances for Blood Collection From Individuals With Hereditary Hemochromatosis;" Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a document entitled "Guidance for Industry: Variances for Blood Collection From Individuals With Hereditary Hemochromatosis" dated August 2001. The guidance document provides recommendations to blood establishments that wish to distribute

blood and blood components collected from individuals with diagnosed hereditary hemochromatosis without indicating the donor's disease on the container label, or collect blood more frequently from such individuals than every 8 weeks without a physical examination and certification of the donor's health by a physician on the day of donation. This guidance document identifies conditions under which FDA will consider approving the above as alternative procedures, or variances, to the current regulations, and provides guidance on what to submit when requesting these variances. These recommendations apply to all blood establishments, whether or not they hold a U.S. license for the manufacture of blood and blood components. The guidance document announced in this notice finalizes the draft guidance document entitled "Guidance for Industry: Variances for Blood Collection From Individuals With Hereditary Hemochromatosis" dated December 2000.

DATES: Submit written or electronic comments on agency guidances at any time.

ADDRESSES: Submit written requests for single copies of this guidance to the Office of Communication, Training, and Manufacturers Assistance (HFM-40), Center for Biologics Evaluation and Research (CBER), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852-1448. Send one self-addressed adhesive label to assist the office in processing your requests. The document may also be obtained by mail by calling the CBER Voice Information System at 1-800-835-4709 or 301-827-1800, or by fax by calling the FAX Information System at 1-888-CBER-FAX or 301-827-3844. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance

Submit written comments on the guidance document to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments.

FOR FURTHER INFORMATION CONTACT: Paula S. McKeever, Center for Biologics Evaluation and Research (HFM–17), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852– 1448, 301–827–6210.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a document entitled "Guidance for Industry: Variances for Blood Collection From Individuals With Hereditary Hemochromatosis" dated August 2001. This guidance document identifies conditions under which FDA will consider approving the above as alternative procedures, or variances, to the current regulations, under the provisions of 21 CFR 640.120 and provides guidance on what to submit when requesting these variances.

On April 29, 1999, the Public Health Service Advisory Committee on Blood Safety and Availability (ACBSA) recommended that the Department of Health and Human Services (DHHS) "create policies that eliminate incentives to seek [blood] donation for purposes of phlebotomy" from patients with diagnosed hemochromatosis who require phlebotomy as therapy for their disease. Further, as undue incentives to donate blood for transfusion (rather than being therapeutically phlebotomized) are removed, DHHS "should create policies that eliminate barriers to using this resource" to augment the country's blood supply (Ref. 1).

On August 10, 1999, the Commissioner of Food and Drugs made a commitment to consider case-by-case exemptions to existing blood labeling and donor suitability regulations for blood establishments that can verify that therapeutic phlebotomy for hemachromatosis is performed at no expense to the patient (Ref. 2). FDA additionally committed itself to work with the Health Care Financing Administration in ensuring that the financial incentives for persons with hereditary hemochromatosis (HH) to donate blood for transfusion are removed. This issue was further discussed at the FDA Blood Products Advisory Committee meeting on September 16, 1999 (Ref. 3). For the foreseeable future, if blood establishments wish to distribute blood collected from donors with HH without disease labeling, they would be responsible for removing financial incentives for these donors. Each blood center should evaluate the advantages of entering these donors into their donor pool.

The guidance document announced in this notice finalizes the draft guidance document entitled "Guidance for Industry: Variances for Blood Collection from Individuals with Hereditary Hemochromatosis" dated December 2000. This guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115; 65 FR 56468, September 19, 2000). This guidance document represents the agency's current thinking on blood collection from individuals with hereditary hemochromatosis. It

does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirement of the applicable statutes and regulations.

II. References

The following references have been placed on display in the Dockets Management Branch (address above) and may be seen by interested persons between 9 a.m. and 4 p.m., Monday through Friday.

- 1. Nightingale, S. D., Summary of Advisory Committee Meeting of April 29 and 30, 1999, May 13, 1999 (http://www.hhs.gov/ bloodsafety).
- 2. Henney, J. E., Memorandum Blood Donations by Individuals with Hemochromatosis, August 1999.
- 3. Blood Products Advisory Committee, 64th Meeting, September 16, 1999 (http://www.fda.gov/ohrms/dockets/ac/acmenu.htm).

III. Comments

Interested persons may, at any time, submit written or electronic comments to the Dockets Management Branch (address above) regarding this guidance document. Two copies of any comments are to be submitted, except individuals may submit one copy. Comments should be identified with the docket number found in the brackets in the heading of this document. A copy of the document and received comments are available for public examination in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

IV. Electronic Access

Persons with access to the Internet may obtain the document at either http://www.fda.gov/cber/guidelines.htm or http://www.fda.gov/ohrms/dockets/ default.htm.

Dated: August 13, 2001.

Margaret M. Dotzel,

Associate Commissioner for Policy. [FR Doc. 01–21247 Filed 8–22–01; 8:45 am] BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of Inspector General

Program Exclusions: July 2001

AGENCY: Office of Inspector General,

ACTION: Notice of program exclusions.

During the month of July 2001, the HHS Office of Inspector General

imposed exclusions in the cases set forth below. When an exclusion is imposed, no program payment is made to anyone for any items or services (other than an emergency item or service not provided in a hospital emergency room) furnished, ordered or prescribed by an excluded party under the Medicare, Medicaid, and all Federal Health Care programs. In addition, no program payment is made to any business or facility, e.g., a hospital, that submits bills for payment for items or services provided by an excluded party. Program beneficiaries remain free to decide for themselves whether they will continue to use the services of an excluded party even though no program payments will be made for items and services provided by that excluded party. The exclusions have national effect and also apply to all Executive Branch procurement and nonprocurement programs and activities.

Subject, city, state Effective date

PROGRAM-RELATED CONVICTIONS

TROOKAM-KEEATED CON	710110140	
ANGELO, ROBERT ANTHONY		(
JR CAMP HILL, PA	08/20/2001	
ANGELO, LILLIAN JAN	08/20/2001	(
BLOOMING GROVE, PA ANSHELL, JACKIE MARLA	08/20/2001	F
DANIA BEACH, FL	00/00/0004	٠
ASKEW, DIANA JOELLHUGOTON, KS	08/20/2001	F
BATIS, DENNIS LYNN LA MESA, CA	08/20/2001	F
BATTLE, STACY ANN	08/20/2001	F
LEAWOOD, KS BECK, JAMES BRAD	08/20/2001	F
HENDERSON, NV BEDROSIAN, YEPRAM	08/20/2001	
CORONA, CA	00/20/2001	F
BLANKENSHIP, KENNETH JOSEPH	08/20/2001	F
LOMPOC, CA	00,20,200.	5
BLANKENSHIP, RUTH CHRIS- TINE EL	08/20/2001	5
SPRING VALLEY, CA BUFALINO, RUSSELL C	08/20/2001	9
CLEARWATER, FL		
CHARNETSKI, KATHYBRYAN, TX	08/20/2001	5
CHARNETSKI, STANLEY HUMBLE, TX	08/20/2001	5
CHEN, YUAN FEI	08/20/2001	5
DIAMOND BAR, CA CONE, ALAN C	08/20/2001	
LOMPOC, CA		5
COOK, CAROL JEAN BLYTHEVILLE, AR	08/20/2001	9
DOLLAR, ZELDER DAVISBORO, GA	08/20/2001	5
ECKERT, RONALD CHARLES	08/20/2001	
OKEMOS, MI GAINES, KENNETH CHARLES	08/20/2001	5
DETROIT, MI GAZES, SHERI L		5
STERLING, VA	08/20/2001	5
HOWELL, NORMAN	08/20/2001	

Subject, city, state	Effective date
BROOKSVILLE, FL KETSOYAN, TIGRAN PASADENA, CA	08/20/2001
KLUDING, CHRISTOPHER SCOTT FORREST CITY, AR	08/20/2001
LAKE, QUANTINA SHALANE S BEND, IN	08/20/2001
LESLIE, FRED LSEMINOLE, FL	08/20/2001
MARTINEZ, TERRY KIRTLAND, NM	08/20/2001
MCNULTY, DENISE MARIE CANBY, OR	08/20/2001
MILLER, EDWIN R BALTIMORE, MD	08/20/2001
MILMAN, BORIS EDISON, NJ	08/20/2001
MIRANDA, ARA RODRIGUEZ DANBURY, CT	08/20/2001
MKROYAN, DAVIS LOS ANGELES, CA	08/20/2001
NEELEY, TERRANCE L OSHKOSH, WI	08/20/2001
NEWLANDER-WINSOR, KIMBERLIN	08/20/2001
ALBUQUERQUE, NM NUTCRACKER BUSINESS	
SVCS, INCLA MESA, CA	08/20/2001
O'DONNELL, JOHN RAY- MOND	08/20/2001
BROOKLYN, NY OVSEPYAN, ZARUHI	08/20/2001
PASADENA, CA PARKS, JAMES DARRELL	08/20/2001
DETROIT, MI PATEL, BIPIN NEW PORT RICHEY, FL	08/20/2001
PRINCE, DAWNMARIE	08/20/2001
RIVERO, CLARA	08/20/2001
RODRIGUEZ, JOSE MIAMI, FL	08/20/2001
ROSEN, NANCY A MCLEAN, VA	08/20/2001
RUKSE, JOSEPH M JR MORGANTOWN, WV	08/20/2001
SCHWARTZ, JEFFREYATLANTIC BEACH, NY	08/20/2001
SCURA, RICHARD JASON LOS ANGELES, CA	08/20/2001
SELBY, JUDITH ANNBEMIDJI. MN	08/20/2001
SELBY, TERRY LEE BEMIDJI, MN	08/20/2001
SEY, SAVON LOS ANGELES, CA	08/20/2001
SHARP, CHRISTOPHER BLAIR AMARILLO, TX	08/20/2001
SHUSTERMAN, SIMON OTISVILLE, NY	08/20/2001
SMITH, ALICE BRYAN, TX	08/20/2001
SORENSON, ROBERT MARCO ISLAND, FL	08/20/2001
STEVENS, ANTHONY ELI ORANGEVALE, CA	08/20/2001
STEVENS, NICKORANGEVALE, CA	08/20/2001
STRANGE, JEFFREY HAYNES	08/20/2001