

sales of bearings in North America, including the brand names under which T&N has sold bearings, must be included in the divestiture.

The proposed Order also addresses a relationship that T&N's thinwall bearings business had with Daido Metals ("Daido"), a Japanese bearing producer. For a number of years, T&N had cooperative technology exchange arrangements with Daido, as well as a joint venture to produce bearings at Bellefontaine, Ohio. In the past, these arrangements between T&N and Daido may have allowed the two companies together to compete better against other bearings producers and to meet their customers' needs for high quality, low cost, sophisticated bearings, better than either company could on its own. To allow for the continuation of cooperation between Daido and the divested T&N bearings business, the proposed Order prohibits Federal-Mogul from entering into such arrangements with Daido for a period of five years. In addition, because certain individuals at T&N are believed to be important to maintaining the cooperative relationships between T&N and Daido, these individuals are given incentives under the proposed Order to stay with the divested T&N thinwall bearings business. The purpose of these provisions is not to force the divested T&N thinwall bearing business or Daido to form any particular cooperative arrangements, but to allow any efficient cooperation between the two firms to continue as if T&N had not been acquired by Federal-Mogul.

The proposed Order also identifies certain assets related to dry bearings or polymer bearings that are to be included in the divestiture. Dry or polymer bearings are bearings that do not rely on a film of oil, but instead on a polymer coating, to reduce friction. These bearings are produced at T&N plants that also produce thinwall bearings, and the inclusion of these bearings in the assets to be divested may be important to the viability of the T&N plants to be divested. Absent the specific references to polymer bearings, the identification of the plants to be divested would require the divestiture of the manufacturing lines for these dry or polymer bearings that are contained in the named plants. However, Federal-Mogul wishes to include these products by name in the proposed Order, to insure the German Federal Cartel Office that the dry bearing products listed will be divested. The German Federal Cartel Office has raised concerns about a product overlap between Federal-Mogul and T&N in dry bearings that would adversely impact competition in dry

bearings in Germany. By including these products in the Commission's proposed Order, Federal-Mogul avoids having to enter into a separate divestiture procedure, relating to the same plants, to satisfy the Federal Cartel Office.

The proposed Order requires that Federal-Mogul divest the identified assets within six months after the proposed Order becomes final. If Federal-Mogul does not divest the assets within that time period, the proposed Order provides for the appointment of a trustee to divest the assets.

The purpose of this analysis is to facilitate public comment on the proposed Order. This analysis is not intended to constitute an official interpretation of the Agreement or the proposed Order or in any way to modify the terms of the Agreement or the proposed Order.

By direction of the Commission, Commissioner Azcuenaga not participating.

Donald S. Clark,

Secretary.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Cardiovascular and Renal Drugs 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: Cardiovascular and Renal Drugs Advisory Committee.

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

Date and Time: The meeting will be held on April 9, 1998, 8:30 a.m. to 5:30 p.m., and April 10, 1998, 8:30 a.m. to 4 p.m.

Location: National Institutes of Health, Clinical Center, Bldg. 10, Jack Masur Auditorium, 9000 Rockville Pike, Bethesda, MD. Parking in the Clinical Center is reserved for Clinical Center patients and their visitors. If you must drive, please use an outlying lot such as Lot 41B. Free shuttle bus service is provided from Lot 41B to the Clinical Center every 8 minutes during rush hour and every 15 minutes at other times.

Contact Person: Joan C. Standaert, Center for Drug Evaluation and Research

(HFD-110), 419-259-6211, or Danyiel D'Antonio (HFD-21), 301-443-5455, Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area), code 12533. Please call the Information Line for up-to-date information on this meeting.

Agenda: On April 9, 1998, the committee will discuss nitric oxide. On April 10, 1998, the committee will discuss new drug applications 20-912 and 20-913, Aggrastat® (tirofiban HCl), Merck Research Laboratories, to be indicated: (1) In combination with heparin for patients with unstable angina or non-Q-wave myocardial infarction to prevent cardiac ischemic events, and (2) patients with coronary ischemic syndromes undergoing percutaneous transluminal coronary angioplasty or atherectomy to prevent cardiac ischemic complications related to abrupt closure of the treated coronary artery.

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 April 2, 1998. Oral presentations from the public will be scheduled between approximately 8:30 a.m. and 9:30 a.m. on April 9, 1998. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before April 2, 1998, and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: March 11, 1998.

Michael A. Friedman,

Deputy Commissioner for Operations.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Peripheral and Central Nervous System Drugs Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.