amendments thereto, is hereby withdrawn, effective December 27, 2005.

In a final rule published elsewhere in this issue of the **Federal Register**, FDA is amending the animal drug regulations to reflect the withdrawal of approval of these NADAs.

Dated: December 7, 2005.

#### Stephen F. Sundlof,

Director, Center for Veterinary Medicine.
[FR Doc. 05–24103 Filed 12–15–05; 8:45 am]
BILLING CODE 4160–01–8

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# **Food and Drug Administration**

Joint Meeting of the Nonprescription Drugs Advisory Committee and the Endocrinologic and Metabolic 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 Committees:
Nonprescription Drugs Advisory
Committee (NDAC) and the
Endocrinologic and Metabolic Drugs
Advisory Committee (EMDAC).

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

Date and Time: The meeting will be held on January 23, 2006, from 8 a.m. to 5 p.m.

Location: Holiday Inn Select Bethesda, Versailles Ballrooms, 8120 Wisconsin Ave., Bethesda, MD. The hotel telephone number is 301–652– 2000

Contact Person: Darrell Lyons, Center for Drug Evaluation and Research (HFD–21), Food and Drug Administration, 5600 Fishers Lane (for express delivery, 5630 Fishers Lane, rm. 1093), Rockville, MD 20857, 301–827–7001, FAX: 301–827–6776, e-mail: lyonsd@cder.fda.gov, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area) codes 3014512541 or 3014512536. Please call the Information Line for upto-date information on this meeting.

Agenda: The committees will consider the safety and efficacy of new drug application (NDA) 21–887, proposing over-the-counter (OTC) use of

ORLISTAT (tetrahydrolipstatin) capsules (60 milligrams (mg)), GlaxoSmithKline Consumer Healthcare, L.P., to promote weight loss in overweight adults when used along with a reduced calorie and low fat diet. The background material will become available no later than the day before the meeting and will be posted under NDAC or EMDAC's docket site at http://www.fda.gov/ohrms/dockets/ac/acmenu.htm (click on the year 2006 and scroll down to NDAC or EMDAC).

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committees. Written submissions may be made to the contact person by January 13, 2006. Oral presentations from the public will be scheduled between approximately 1 p.m. and 2 p.m. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person before January 13, 2006, 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.

Persons attending FDA's advisory committee meetings are advised that the agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Darrell Lyons at least 7 days in advance of the meeting.

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

Dated: December 2, 2005.

#### Jason Brodsky,

Acting Associate Commissioner for External Relations.

[FR Doc. 05–24101 Filed 12–15–05; 8:45 am] BILLING CODE 4160–01–S

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [FDA 225-05-8000]

Memorandum of Understanding Between the United States Food and Drug Administration and the C-Path Institute

**AGENCY:** Food and Drug Administration,

HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is providing notice of a memorandum of understanding (MOU) between the United States Food and Drug Administration and the C-Path Institute. The specific purpose of this MOU is to establish an overarching framework for collaboration between the parties. This framework will be based on mutually agreed upon programs and activities in the areas of applied scientific research and training/ education to foster the development of new evaluation tools to inform medical product development. The parties shall each leverage its own expertise and resources to facilitate programs of shared interests across the diverse disciplines of therapeutics, biological sciences, engineering and medical devices in building applied research and training/education programs. The appropriate formal agreements will be executed as required by law for any activities that result from this collaboration.

**DATES:** The agreement became effective October 14, 2005.

FOR FURTHER INFORMATION CONTACT: For C-Path Institute: Raymond L. Woosley, The Critical Path Institute, 4280 N. Campbell Ave., #214, Tucson, AZ 85718, 520–547–3440, FAX: 520–547–3456, e-mail: rwoosley@c-path.org.

For The Food and Drug Administration: Mary I. Poos, Office of External Relations, Food and Drug Administration (HF–10), 5600 Fishers Lane, Rockville, MD 20857, 301–827–2825, FAX: 301–827–3042, e-mail: mary.poos@fda.gov.

SUPPLEMENTARY INFORMATION: In accordance with 21 CFR 20.108(c), which states that all written agreements and MOUs between FDA and others shall be published in the **Federal** Register, the agency is publishing notice of this MOU.

Dated: December 7, 2005.

#### Jeffrey Shuren,

Assistant Commissioner for Policy.

BILLING CODE 4160-01-S

225-05-8000

# MEMORANDUM OF UNDERSTANDING BETWEEN THE UNITED STATES FOOD AND DRUG ADMINISTRATION AND THE C-PATH INSTITUTE

This Memorandum of Understanding (MOU) between the U.S. Food and Drug Administration (FDA) and the Critical Path Institute (C-Path) (hereafter termed "the Parties") formalizes an agreement between the two parties to develop collaborative activities in the areas of applied research, training and education to enhance safe and efficacious medical product development.

## I. Purpose

The specific purpose of this MOU is to establish an overarching framework for collaboration between the Parties. This framework will be based on mutually agreed upon programs and activities in the areas of applied scientific research and training/education to foster the development of new evaluation tools to inform medical product development. The Parties shall each leverage its own expertise and resources to facilitate programs of shared interests across the diverse disciplines of therapeutics, biological sciences, engineering and medical devices in building applied research and training/education programs. The appropriate formal agreements will be executed as required by law for any activities that result from this collaboration.

# II. Background

The FDA is responsible for reviewing clinical research to ensure that marketed human medical products (drugs, biologics, and medical devices) have been shown to be safe and effective.

The C-Path Institute is a non-profit research and education organization located in Tucson, Arizona. The Institute's purpose is to create innovative programs in education and research to enable safe acceleration of medical product development. It also serves as a 'neutral ground' for academia, industry and government to test ideas that will result in optimal (safe, effective, timely) drug development processes. C-Path brings together faculty from the UAZ Colleges of Pharmacy, Medicine, Agriculture and Life Sciences, and the School of Management as well as clinicians and researchers from the UAZ Comprehensive Cancer Center, the Sarver Heart Center, the Pima Community College, Arizona State University and the Translational Genomics Research Institute in programs related to pharmaceutical discovery and development, clinical research and good clinical practices (GCP) as well as scientific staff from SRI International (an independent, non-profit technology development organization), who have substantial experience in developing drugs for commercial manufacturing. SRI International is a contractor for NIH and has initiated a drug development consortium with other academic institutions, the purpose of which is to assist faculty investigators to translate research into clinical drug candidates.

# III. Substance of Agreement

This MOU is intended as an overarching framework for joint collaboration between the Parties, toward the goal of developing new evaluative tools to inform medical product development. The areas of collaboration would include, but not be limited to:

<u>Training/Education programs</u>: Activities arising from complementary interests will be developed jointly by C-Path and FDA, and offered to academia, industry, and others as identified needs arise. The Parties will disseminate information through mutually agreed vehicles including training activities, meetings, and symposia.

<u>Applied Research programs:</u> Programs will be developed in areas of mutual complementary interest such as imaging, biomarkers and surrogate markers, proteomics and genomics, clinical trial design, and other areas that will enhance medical product development.

As specific topics for joint training/education and/or research are identified under this MOU they will be conducted under the appropriate formal agreements as required by law.

# IV. Participation

It is anticipated that a wide range of faculty and graduate students, clinicians, and researchers from academic programs may participate in activities developed under this agreement, including, but not limited to, University of Arizona Comprehensive Cancer Care Center, the Sarver Heart Center, the Colleges of Pharmacy, Medicine, Management, and Agriculture and Life Sciences, Pima Community College, Arizona State University, Translational Genomics Research Institute, and SRI International. Other participants could include FDA staff, scientists from industry, field laboratories and others identified for joint training and outreach activities.

Each Party will appoint appropriate representatives to facilitate the planning, preparation, and implementation of the activities within the framework of this MOU. Meetings will be convened at a venue and time agreed between Parties, and each Party shall be responsible for its own expenses incurred in sending representatives to these meetings.

# V. Resource Obligations

This MOU describes in general terms the basis upon which the Parties intend to cooperate. It does not create binding, enforceable obligations against any Party. All activities undertaken pursuant to the MOU are subject to the availability of personnel, resources, and appropriated funds. This MOU does not affect or supersede any existing or future agreements or arrangements among the Parties and does not affect the ability of the Parties to enter into other agreements or arrangements related to this MOU.

# VI. Name and Address of Participating Parties and Liaisons

#### A. C-Path Institute

Raymond L. Woosley, M.D., Ph.D.

President

The Critical Path Institute 4280 N. Campbell Ave. # 214

Tucson, AZ 85718 Phone: 520-547-3440 Fax: 520-547-3456

Email: rwoosley@c-path.org

# B. Food and Drug Administration

Mary I. Poos, Ph.D.

Director, Academic and Intellectual Partnerships

Office of External Relations
Food and Drug Administration

Parklawn Bldg, Room 14C-06 (HF-10)

5600 Fishers Lane Rockville, MD 20857

Tel: (301) 827-2825 Fax: (301) 827-3042

Email: mary.poos@fda.gov

# VII. Period of Agreement

This MOU becomes effective upon the date of the last Party to sign ("effective date") and will continue in effect for five years. It may be modified by mutual written consent or terminated by either party upon a 30-day advanced written notice to the other party. The Parties agree to evaluate the MOU periodically during the effective period, but at least once annually, on or before the anniversary of the effective date. Upon evaluation, either Party shall have the option of continuing, modifying, or canceling this agreement as provided for in Article VII of this MOU.

APPR	OVED	AND	<b>ACCEPTED</b>	<b>FOR</b>	THE

C-PATHANSTITUTE

APPROVED AND ACCEPTED FOR THE FOOD AND DRUG ADMINISTRATION

Title Deputy Commissioner for Operations

Food and Drug Administration

Cleared: R. Garwood, ORM 5/18/05

Reviewed and cleared: R. Springer, OM/OAGS 5/18/05

Reviewed and edited: L. Mahler, OCC 8/23/05 Reviewed and cleared: P. Stannard, OS 9/12/05

By (

[FR Doc. 05-24100 Filed 12-15-05; 8:45 am] BILLING CODE 4160-01-C

### **DEPARTMENT OF HEALTH AND HUMAN SERVICES**

### Office of Inspector General

### **Program Exclusions: November 2005**

**AGENCY:** Office of Inspector General, HHS.

**ACTION:** Notice of program exclusions.

During the month of November 2005, the HHS Office of Inspector General imposed exclusions in the cases set forth below. When an exclusions 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 non-

proguroment programs and a	otivitios	SUNAEFEN, UNNISTA	12/20/2005		
procurement programs and activities.		MARICOPA, AZ TAMAYO, HEIROL	12/20/2005	ANDERSON, ERNEIS	12/20/2005
0.11	Effective		12/20/2005	CLINTON, SC	
Subject name, address	date	MONTGOMERY, AL	40/00/0005	BASSO, ALICIA	12/20/2005
		VALLE, LUIS	12/20/2005	ROCHESTER, NY	
PROGRAM-RELATED CONVICTIONS		GLENDALE, CA		CANTU, HELEN	12/20/2005
- THOGHAM NELATED GOIN	710110110	VUKASIN, ALAN	12/20/2005	MADERA. CA	, ,
BASONE, RENNAE	12/20/2005	COTTONWOOD, ID		CLARK, RUBY	12/20/2005
AKRON, OH	12/20/2000	WEILAND, JEANETTE	12/20/2005	PIONEER. LA	12/20/2000
BERTIE, LIONEL	12/20/2005	HURSON, SD		CLENDENEN, BRENDA	12/20/2005
	12/20/2005	WHITE, ROBERT	12/20/2005		12/20/2003
TOLEDO, OH	10/00/0005	WILSONVILLE, OR		HODGEN, OK	10/00/0005
BRAVO, THERESA	12/20/2005			COPES, RESHAWN	12/20/2005
SAN DIEGO, CA		FELONY CONVICTION FOR HE	ALTH CARE	LEESVILLE, LA	
BROUSSARD, JERRY	12/20/2005	FRAUD		HAWKINS, BRIAN	12/20/2005
BEAUMONT, TX				FOUNTAIN HILLS, AZ	
BRUMBAUGH, JAY	12/20/2005	BUNKER, JASON	12/20/2005	JIMENEZ, ALICIA	12/20/2005
COLLINSVILLE, OK		SAN BERNARDINO, CA		LOS ANGELES, CA	
CABRERA, DAISY	12/20/2005	CATANZARO, DANIEL	12/20/2005	LAMBERT, STEPHANIE	12/20/2005
BRONX, NY		CARTERSVILLE, GA		JANESVILLE, WI	
CARTER, ANGELA	12/20/2005	DENNETT, ROBIN	12/20/2005	MOENICH, KIM	12/20/2005
OSKALOOSA, IA		AUGUSTA. ME		CLEVELAND, OH	
CHINI, JERI	12/20/2005	HAMPTON, STACEY	12/20/2005	MOORE, MICHAEL	12/20/2005
PORT CLINTON, OH		ST LOUIS. MO		HENNESSEY, OK	
CLOSE, CHRISTOPHER	12/20/2005	MOORE, MARK	12/20/2005	NAVARRO, JEA	12/20/2005
ATWATER, CA		DAYTON, OH		VALLEJO, CA	
DYE, HEATHER	12/20/2005		12/20/2005	NORRIS, KIMBERLY	12/20/2005
MILWAUKEE. WI		OKLAHOMA CITY, OK		CHILDRESS, TX	
EDWARDS, TERRI	12/20/2005		12/20/2005	POPPY, THOMAS	12/20/2005
GAHANNA, OH		COLUMBUS. OH	, _ 0, _ 0 0	SPRING HILL, FL	,_,,_,
FRID, BORIS	12/20/2005		12/20/2005	PRADA, GERMAN	12/20/2005
WOODLAND HILLS, CA	. =, = 3, = 000	SANTA CLARA, CA	12,23,2000	SPRINGBORO. OH	. =, = 3, = 000
	12/20/2005	WHITE, JACQUESE	12/20/2005		12/20/2005
GG: 1221011, G10/1111	.2,20,2000		.2,20,2000	57.555.55, VEITONION	. 2, 20, 2000

Subject name, address	Effective date	Subject name, address	Effective date
BLOOMFIELD HILLS, MI		MEMPHIS, TN	
HARVEY, JAMES	12/20/2005		
LAKEWOOD, WA		FELONY CONTROL SUBS	STANCE
HARVEY, RUBY	12/20/2005	CONVICTION	
LAKEWOOD, WA		011111071	
HATHAWAY, BRIAN	12/20/2005	CHLYSTA, RUSSELL	12/20/2005
REYNOLDSBURG, OH	40/00/0005	YANKTON, SD	10/00/0005
HOWARD, JULIE	12/20/2005	FERNANDEX, BRENDA	12/20/2005
FULTON, MS KHANNA, ARUN	12/20/2005	NACOGDOCHES, TX GRAYS, SONYA	12/20/2005
STONE MOUNTAIN, GA	12/20/2005	WACO, TX	12/20/2005
LOCKE, STEPHANIE	12/20/2005	HAAKE, DONNA	12/20/2005
HOUSTON, TX	12/20/2003	BELLEVUE, NE	12/20/2003
MASON, CLINT	12/20/2005	HEIKENS, ANGELA	12/20/2005
FORK, SC	12/20/2003	BELLE FOURCHE, SD	12/20/2003
MERRITT, RICKLEY	12/20/2005	KUTZNER, JAMES	12/20/2005
GREER, SC	, ,	LOUISVILLE, KY	12/20/2000
MILLER, MICHELLE	12/20/2005	MARTENS, DALE	12/20/2005
SHAKOPEE, MN		LONDONDERRY, VT	12/20/2000
MOORE, MERLYN	12/20/2005	MCCARTNEY, LUANNE	12/20/2005
RIALTO, CA		COAHOMA, TX	
MR J'S LIQUOR, INC	12/20/2005	NYMAN, CATHERINE	12/20/2005
LOS ANGELES, CA		DENVER, CO	
PABBATHI, RAMMOHAN	12/20/2005	ODVODY, DAWN	12/20/2005
TRENTON, NJ		GREENVILLE, IL	
PROFESSIONAL AMBU-		PALMER, MARTIN	12/20/2005
LANCE SVC OF NORWICH,		WASILLA, AK	
INC	12/20/2005	RIOJAS, JEANETTE	12/20/2005
NORWICH, CT		DEER PARK, TX	
PROVINCE, KIMBERLY	12/20/2005	RYABIK, BRETT	12/20/2005
CLARKSDALE, MS	10/00/0005	DORAVILLE, GA	
RINGGENBERG, JULIE	12/20/2005	SCOTT, BRUCE	12/20/2005
OTTUMWA, IA ROBY, JARROD	10/00/0005	QUINCY, IL	
COLUMBUS, OH	12/20/2005	SHULTZ, ALAN	12/20/2005
ROSEL, NICOLE	12/20/2005	MOUNT STERLING, KY	
COOPERSVILLE, MI	12/20/2005	WAY, NANCY	12/20/2005
SATTARI, PARI	12/20/2005	FT WORTH, TX	
TARZANA, CA	12/20/2003	WHITAKER, DARWIN	12/20/2005
SAULTER, MONROE	12/20/2005	HAZARD, KY	
THREE RIVERS, TX	12/20/2000	DATIENT ADVICE/NECL FOT C	211//27/21/2
SCHAEFER, CHRISTA	12/20/2005	PATIENT ABUSE/NEGLECT CO	DNVICTIONS
MARICOPA, AZ		ANDERCON EDNELO	10/00/0005
TAMAYO, HEIROL	12/20/2005	ANDERSON, ERNEIS	12/20/2005
MONTGOMERY, AL		CLINTON, SC BASSO, ALICIA	10/00/0005
VALLE, LUIS	12/20/2005	ROCHESTER. NY	12/20/2005
GLENDALE, CA		CANTU, HELEN	12/20/2005
VUKASIN, ALAN	12/20/2005	MADERA. CA	12/20/2005
COTTONWOOD, ID		CLARK. RUBY	12/20/2005
VALCH AND IDANIETTE	10/00/000	OFVUL, UODI	1 12/20/2003