

and discussed at this meeting. These discussions are likely to reveal personal information concerning individuals associated with the applications. This information is exempt from mandatory disclosure under the above-cited statutes.

*SEP Meeting on:* AHRQ Limited Competition: PROSPECT STUDIES—Building New Clinical Infrastructure for CE (R01).

*Date:* June 16, 2010 (Open on June 16 from 8 a.m. to 8:15 a.m. and closed for the remainder of the meeting).

*Place:* Hyatt Regency Bethesda Hotel, 7400 Wisconsin Avenue, 1 Bethesda Metro Center, Bethesda, Maryland 20814.

*Contact Person:* Anyone wishing to obtain a roster of members, agenda or minutes of the non-confidential portions of this meeting should contact Mrs. Bonnie Campbell, Committee Management Officer, Office of Extramural Research, Education and Priority Populations, AHRQ, 540 Gaither Road, Room 2038, Rockville, Maryland 20850, Telephone (301) 427-1554.

Agenda items for this meeting are subject to change as priorities dictate.

Dated: May 24, 2010.

**Carol M. Clancy,**  
*Director.*

[FR Doc. 2010-13109 Filed 6-2-10; 8:45 am]

**BILLING CODE 4160-90-M**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Agency for Healthcare Research and Quality; Notice of Meeting

In accordance with section 10(d) of the Federal Advisory Committee Act (5 U.S.C., Appendix 2), announcement is made of a Health Care Policy and Research Special Emphasis Panel (SEP) meeting.

A Special Emphasis Panel is a group of experts in fields related to health care research who are invited by the Agency for Healthcare Research and Quality (AHRQ), and agree to be available, to

conduct on an as-needed basis, scientific reviews of applications for AHRQ support. Individual members of the Panel do not attend regularly-scheduled meetings and do not serve for fixed terms or a long period of time. Rather, they are asked to participate in particular review meetings which require their type of expertise.

Substantial segments of the upcoming SEP meeting listed below will be closed to the public in accordance with the Federal Advisory Committee Act, section 10(d) of 5 U.S.C., Appendix 2 and 5 U.S.C. 552b(c)(6). Grant applications for the OS ARRA: Optimizing Prevention and Healthcare Management for Complex Patients (R21) applications are to be reviewed and discussed at this meeting. These discussions are likely to reveal personal information concerning individuals associated with the applications. This information is exempt from mandatory disclosure under the above-cited statutes.

*SEP Meeting on:* OS ARRA: Optimizing Prevention and Healthcare Management for Complex Patients (R21).

*Date:* June 24, 2010 (Open on June 24 from 8 a.m. to 8:15 a.m. and closed for the remainder of the meeting).

*Place:* Hilton Rockville Executive Meeting Center, 1750 Rockville Pike, Conference Room TBD, Rockville, MD 20850.

*Contact Person:* Anyone wishing to obtain a roster of members, agenda or minutes of the nonconfidential portions of this meeting should contact Mrs. Bonnie Campbell, Committee Management Officer, Office of Extramural Research, Education and Priority Populations, AHRQ, 540 Gaither Road, Room 2038, Rockville, Maryland 20850, Telephone (301) 427-1554.

Agenda items for this meeting are subject to change as priorities dictate.

Dated: May 24, 2010.

**Carolyn M. Clancy,**  
*Director.*

[FR Doc. 2010-13108 Filed 6-2-10; 8:45 am]

**BILLING CODE 4160-90-M**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA-2010-N-0004]

[FDA 225-09-0014]

### Memorandum of Understanding by and Between the United States Food and Drug Administration and the International Anesthesia Research Society for the Safety of Key Inhaled and Intravenous Drugs in Pediatrics Public-Private Partnership

**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 FDA and the International Anesthesia Research Society (IARS). The purpose of this MOU is to establish a framework for collaboration between FDA and IARS and to support their shared interest of promoting the safe use of anesthetics and sedatives in children.

**DATES:** The agreement became effective March 21, 2010.

**FOR FURTHER INFORMATION CONTACT:** Wendy R. Sanhai, Senior Scientific Advisor, Office of the Commissioner, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, rm. 4128, Silver Spring, MD 20993, 301-796-8518.

**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: May 26, 2010.

**Leslie Kux,**

*Acting Assistant Commissioner for Policy.*

**BILLING CODE 4160-01-S**

225-09-0014

**MEMORANDUM OF UNDERSTANDING  
BY AND BETWEEN THE**

**UNITED STATES FOOD AND DRUG ADMINISTRATION (FDA)**

**AND THE**

**INTERNATIONAL ANESTHESIA RESEARCH SOCIETY (IARS)**

**FOR**

**THE SAFETY OF KEY INHALED AND INTRAVENOUS DRUGS IN PEDIATRICS**

**PUBLIC-PRIVATE PARTNERSHIP (SAFEKIDS PPP)**

This Memorandum of Understanding (MOU) is executed by and between the United States Food and Drug Administration (FDA) and the International Anesthesia Research Society (IARS), hereafter referred to collectively as the "Parties." This MOU is deemed effective as of the date it is fully signed by both Parties (Effective Date).

WHEREAS, non-clinical studies in juvenile animal models show that exposure to some anesthetics and sedatives is associated with memory and learning deficits and other neurodegenerative changes in the central nervous system;

WHEREAS, insufficient human data exist to support or refute the possibility that similar effects could occur in children; thus, there is a critical need to address the public health issues associated with the safe use of anesthesia and sedatives in children;

WHEREAS, the FDA, under its public health mission, is interested in partnering with multiple stakeholders (e.g. professional societies, academic research institutions, patient advocacy groups, industry and other government and nonprofit organizations) to investigate the effect of anesthetics and sedatives on the developing human brain, including long-term studies in neonates and young children, and to ensure that information and outcomes generated from this research can be used to benefit public health;

WHEREAS, the IARS is a nonpolitical nonprofit voluntary membership society, organized and operated exclusively for exempt purposes as set forth in section 501(c)(3) of the Internal Revenue Code, whose mission is to encourage, stimulate, and fund ongoing anesthesia related research projects and to disseminate current, state-of-the-art, basic and clinical research data in all areas of clinical anesthesia;

WHEREAS, the FDA and the IARS seek to develop a Public-Private Partnership (PPP) to leverage the resources and expertise of the Parties and develop an overarching framework to bring together multiple stakeholders to address the major scientific and clinical gaps regarding the safe use of anesthetics and sedatives in children;

WHEREAS, the Parties have agreed to enter into this MOU to develop the SAFEKIDS (Safety of Key Inhaled and Intravenous Drugs in Pediatrics) PPP, a multi-year, multi-phased collaborative effort to make anesthesia safer for children, which will include multiple public and private partners working together in the interest of public health.

NOW, THEREFORE, in consideration of the mutual agreement of the Parties, and of the covenants and conditions hereinafter expressed, the Parties hereby agree as follows:

## **I. PURPOSE**

The purpose of this MOU is to establish the framework for collaboration between the Parties and to support their shared interest of promoting the safe use of anesthetics and sedatives in children. The strategic goals and expected results of this collaboration are to:

1. **Establish the SAFEKIDS PPP for the purpose of supporting, implementing, and managing a series of scientific projects to bridge the knowledge gaps that exist in elucidating whether certain anesthetic and sedative agents cause neurotoxicity in rodents, non-human primates, and humans.**
2. **Share information and data to the extent permitted by State and Federal law, and ensure that information, know-how, and best practices resulting from the scientific projects conducted under the SAFEKIDS PPP are placed in the public domain for the benefit of all stakeholders.**
3. **Inform clinicians, patients, and other stakeholders about any potential safety risks associated with certain anesthetic and sedative agents, through joint publications, workshops, and other educational efforts.**
4. **Inform future activities, including clinical trials, and the research and development of anesthetic and sedative agents for intended use in the pediatric population, to the extent permitted by available scientific data.**

## **II. AUTHORITY**

**FDA is authorized to enforce the Federal Food, Drug, and Cosmetic Act (the Act) as amended (21 U.S.C. 301). In fulfilling its responsibilities under the Act, FDA among other things, directs its activities toward promoting and protecting the public health by assuring the safety, efficacy, and security of drugs. To accomplish its mission, FDA must stay abreast of the latest developments in research and also communicate with stakeholders about complex scientific and public health issues. Increased development of research, education and outreach partnerships with IARS will greatly contribute to FDA's mission.**

## **III. RESPONSIBILITIES OF THE PARTIES**

**In pursuit of the goals described above, the Parties agree to work through the following process.**

1. **Under the framework of the SAFEKIDS PPP, the FDA and the IARS will develop an overarching infrastructure to implement and sustain additional pre-clinical and clinical research, leveraging a combination of public and private resources and expertise to bridge the scientific and public health gaps in the pediatric population.**
2. **The Parties will establish a public-private governance structure including a Steering Committee, an Executive Board, a Scientific Advisory Board, and technical subcommittees. These governance committees will develop strategic and operational plans, set priorities, and review, implement, oversee, and evaluate individual projects conducted under the SAFEKIDS PPP. The membership of the governance committees will be inclusive and will be comprised of representatives of the Parties and other stakeholders. The Parties will develop specific policies and procedures to carry out the work of the governance committees within the framework of the PPP.**

No committee, board, or subcommittee established pursuant to this MOU will provide advice or recommendations to FDA or to any government agency.

3. To ensure integrity, scientific rigor, and consistency with the goals and objectives of the SAFEKIDS PPP, all projects proposed for implementation will undergo peer review, with final project selection to be accomplished through a consensus among qualified experts operating in a consistent and unbiased manner within the overall governance structure of the PPP.
4. Data, outcomes and best practices generated under the SAFEKIDS PPP will be placed in the public domain for the benefit of all stakeholders and patients.

#### **IV. RESOURCES**

Sources of support for projects under this MOU will be governed by State and Federal law and applicable policies and procedures. The terms for such support will be set forth in the specific and separate written agreements for each project. The MOU 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 funds. This MOU does not affect or supersede any existing or future agreements or arrangements among the Parties. This MOU and all associated agreements will be subject to the applicable policies, rules, regulations, and statutes under which the FDA and IARS operate.

#### **V. GENERAL PROVISIONS**

1. Nothing in this MOU alters the statutory authorities or obligations of FDA. This MOU is intended to facilitate cooperative efforts between the Parties in the area of pediatric anesthesiology.
2. U.S. Federal law governs this MOU for all purposes, including, but not limited to, determining the validity of the MOU, the meaning of its provisions, and the rights, obligations, and remedies of the Parties.
3. Access to non-public information shall be governed by separate Confidentiality Disclosure Agreements in which the Parties will agree and certify in writing that they shall not further release, publish or disclose such information and that they shall protect such information. No proprietary data, trade secrets or patient confidential information shall be disclosed among the Parties unless permitted by the provisions of 21 U.S.C. 331(j), 21 U.S.C. 360j(c), 18 U.S.C. 1905, and other pertinent laws and regulations governing the confidentiality of such information.
4. Release of information to the media or to the general public about the SAFEKIDS PPP or the activities conducted by the Parties pursuant to the PPP and this MOU shall be subject to prior review by and agreement between the Parties.

5. It is understood that, although the Parties have mutual interests, there may be opportunities for independent collaborations and activities outside the scope of this MOU, but which are within the scope of the Parties' respective missions. As such, the Parties may, as appropriate, enter into independent negotiations and agreements with prospective partner/s without any effect on this MOU.
6. Rights to inventions or intellectual property developed will be addressed in separate written development and implementation agreements among the Parties. To the extent there is FDA participation in any projects related to development of any product, invention, or property developed, such activities will be governed by applicable Federal law.
7. Any notice or other communication required or permitted under this MOU shall be in writing and will be deemed effective on the date it is received by the receiving Party.
8. FDA participation in this MOU is governed by Federal statutes and regulations.

## **VI. TERM, TERMINATION AND MODIFICATIONS**

1. This MOU constitutes the entire agreement between the Parties as to the matters herein. There are no representations, warranties, agreements, or understandings, expressed or implied, written or oral, between the Parties relating to the subject matter of this MOU that are not fully expressed herein.
2. This MOU may be modified only upon the mutual written consent of the Parties. Modifications must be signed by the original signatories to this MOU, or by their designees or successors. No oral statement by any person shall be interpreted as modifying or otherwise affecting the terms of this MOU.
3. This MOU, when accepted by the Parties, will remain in effect for three (3) calendar years from the Effective Date, unless modified or terminated.
4. Either Party to this MOU may terminate its participation by written notice at any time, with or without cause, and without incurring any liability or obligation. Such written notice shall be given by the terminating Party to the other Party at least 60 days prior to the date of actual termination.

## **VII. CONTACTS**

Notices or formal communications pursuant to this MOU shall be sent in writing by personal delivery, overnight delivery, facsimile telecommunication with confirmatory receipt, or certified or registered mail, return receipt requested, to the following contact for each Party:

For FDA: Wendy R. Sanhai, Ph.D., M.B.A.  
Senior Scientific Advisor  
Office of the Commissioner, FDA  
5600 Fishers Lane, Suite 6A-08  
Rockville, MD. 20857  
Fax: (301) 827-5891

With a copy to: Chekesha S. Clingman, Ph.D.  
Senior Scientific Program Manager  
Office of the Commissioner, FDA  
5600 Fishers Lane, Suite 6A-08  
Rockville, MD 20857  
Fax: (301) 827-5891

For IARS: Robert N. Sladen, MD  
Chair, IARS Board of Trustees  
International Anesthesia Research Society  
100 Pine Street, Suite 230  
San Francisco, CA 94111  
Fax: (415) 296-6901

With a copy to: Thomas A. Cooper  
Executive Director  
International Anesthesia Research Society  
100 Pine Street, Suite 230  
San Francisco, CA 94111  
Fax: (415) 296-6901

The Parties shall notify each other of any change of address or change of named contact by written notice as specified in this paragraph VI. All notices shall be effective upon date of receipt.

**Signatures begin on next page**

**SIGNATURES OF RESPONSIBLE PARTIES:**

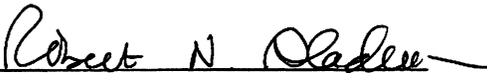
We, the undersigned, agree to abide by the terms and conditions of this MOU.

APPROVED AND ACCEPTED FOR THE  
UNITED STATES FOOD AND DRUG ADMINISTRATION

  
\_\_\_\_\_  
Janet Woodcock, M.D.  
Director, Center for Drug Evaluation and Research  
Food and Drug Administration

Date 3/20/10

APPROVED AND ACCEPTED FOR THE  
INTERNATIONAL ANESTHESIA RESEARCH SOCIETY

  
\_\_\_\_\_  
Robert N. Sladen, M.D.  
Chair, IARS Board of Trustees  
International Anesthesia Research Society

Date 3-21-2010

*FDA/IAR SAFEKIDS PPP*  
*Page 7 of 7*

[FR Doc. 2010-13292 Filed 6-2-10; 8:45 am]  
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**DEPARTMENT OF HEALTH AND  
HUMAN SERVICES****Administration for Children and  
Families****Office of Child Support Enforcement;  
Privacy Act of 1974; Computer  
Matching Agreement**

**AGENCY:** Office of Child Support  
Enforcement (OCSE), ACF, HHS.

**ACTION:** Notice of a Computer Matching  
Program.

**SUMMARY:** In accordance with the  
Privacy Act of 1974 (5 U.S.C. 522a), as  
amended, OCSE is publishing notice of  
a computer matching program between  
OCSE and State agencies administering  
unemployment compensation (UC)  
programs.

**DATES:** The Department of Health and  
Human Services (HHS) invites  
interested parties to review, submit  
written data, comments, or arguments to  
the agency about the matching program  
until July 6, 2010. As required by the  
Privacy Act (5 U.S.C. 552a(r)), HHS on