submitted manuscripts, and grant applications submitted to NIH in 2004, he falsely claimed that transgenic mice had been generated with the monofunctional lentiviral vectors with c-Mvc, Ras or Akt under the control of the CD4 promoter, when they had not, and that transgenic mice had been generated with the bi-functional lentiviral vectors with CD4-c-Myc, Ras or Akt- and U6shRNAs targeting luciferase, Bcl-2, or Bim proteins, when they had not. The effect of these misrepresentations was the reported false conclusion that a cytokine-stimulated proto-oncogene network regulated CD4+ T-cell survival and responses to foreign and self antigens.

8. While at MIT, Dr. Luk Van Parijs admitted that in presentations and submitted manuscripts in 2004, he falsely claimed that mice injected with plasmids carrying shRNAs for Bcl–2, Akt1 and Akt2, complexed to polyethylene imine (PEI) showed a significant reduction in c–myc-induced tumor growth, when the experiments had not been done.

9. While at MIT, Dr. Luk Van Parijs admitted that in presentations in 2004, he falsely claimed that shRNAs designed using algorithms developed in 2004 were more effective to silence target genes than the shRNAs designed with algorithms in 2002.

10. While at MIT, Dr. Luk Van Parijs admitted that in multiple presentations, submitted manuscripts, a grant application submitted to NIH, and in the text of Current Opinions in Molec. Therapeutics, 6:136, 2004, he falsely claimed that an in vivo RNAi screen was developed to identify genes in cytokine and apoptosis pathways that accelerated or suppressed Myc-induced tumorigenesis in lethally irradiated mice, by using bi-functional lentiviral vectors that expressed c-Myc under control of the CMV enhancer-β-actin promoter (CAG) and U6-driven shRNAs designed to silence 168 selected genes, when the experiments had not been done.

11. While at MIT, Dr. Luk Van Parijs admitted that in a submitted manuscript in 2004 and a grant application submitted to NIH in 2003, he falsely claimed that with the use of retroviral vectors with Bim and activated Ras, Akt or Myc, he showed that the IL–2stimulated activation of proto-oncogene pathways functioned to promote the survival of T cells following antigen encounter by regulating Bim and Bcl–2 pathways, when the experiments that were performed were inconclusive.

Dr. Van Parijs has entered into a Voluntary Exclusion Agreement in which he has voluntarily agreed, for a period of five (5) years, beginning on December 22, 2008:

(1) to exclude himself from any contracting or subcontracting with any agency of the United States Government and from eligibility or involvement in nonprocurement programs of the United States Government referred to as "covered transactions" pursuant to HHS' Implementation (2 CFR Part 376 *et seq.*) of OMB Guidelines to Agencies on Government wide Debarment and Suspension (2 CFR, Part 180); and

(2) To exclude himself from serving in any advisory capacity to PHS, including but not limited to service on any PHS advisory committee, board, and/or peer review committee, or as a consultant.

FOR FURTHER INFORMATION CONTACT:

Director, Division of Investigative Oversight, Office of Research Integrity, 1101 Wootton Parkway, Suite 750, Rockville, MD 20852, (240) 453–8800.

Dated: January 14, 2009.

Chris B. Pascal,

Director, Office of Research Integrity. [FR Doc. E9–1453 Filed 1–22–09; 8:45 am] BILLING CODE 4150–31–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Committee on Vital and Health Statistics: Meeting

Pursuant to the Federal Advisory Committee Act, the Department of Health and Human Services (HHS) announces the following advisory committee meeting.

Name: National Committee on Vital and Health Statistics (NCVHS), Full Committee Meeting.

Time and Date:

February 25, 2009, 9 a.m.–3 p.m. February 26, 2009, 10 a.m.–4 p.m.

Place: Hubert Humphrey Building, 200 Independence Avenue, SW., Room 505A, Washington, DC 20201.

Status: Open.

Purpose: At this meeting the Committee will hear presentations and hold discussions on several health data policy topics. On the morning of the first day the Committee will hear updates from the Department, the HHS Data Council, the Center for Medicare and Medicaid Services, as well as update on the transition to the new administration. There will also be an ONC update on the NHIN Conference. In the afternoon there will be a speaker on de-identification of health data from the Center for Democracy and Technology.

On the morning of the second day there will be a briefing on international terminology and an update on Health Statistics for the 21st Century. There will also be an update from NCHS Board of Scientific Counselors and an overview of emerging and innovative sources of health data. The times shown above are for the full Committee meeting. Subcommittee breakout sessions can be scheduled for late in the afternoon of the first day and second day and in the morning prior to the full Committee meeting on the second day. Agendas for these breakout sessions will be posted on the NCVHS website (URL below) when available.

For Further Information Contact: Substantive program information as well as summaries of meetings and a roster of committee members may be obtained from Marjorie S. Greenberg, Executive Secretary, NCVHS, National Center for Health Statistics, Centers for Disease Control and Prevention, 3311 Toledo Road, Room 2402, Hyattsville, Maryland 20782, telephone (301) 458–4245. Information also is available on the NCVHS home page of the HHS Web site: http:// www.ncvhs.hhs.gov/, where further information including an agenda will be posted when available.

Should you require reasonable accommodation, please contact the CDC Office of Equal Employment Opportunity on (301) 458–4EEO (4336) as soon as possible.

Dated: January 12, 2009.

James Scanlon,

Deputy Assistant Secretary for Science and Data Policy, Office of the Assistant Secretary for Planning and Evaluation.

[FR Doc. E9–1445 Filed 1–22–09; 8:45 am] BILLING CODE 4151–05–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-10273]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

1. Type of Information Collection Request: New collection; Title of Information Collection: Evaluation of the Medicare Care Management Performance Demonstration (MCMP) and the Electronic Health Records Demonstration (EHRD); Use: The MCMP demonstration was authorized under Section 649 of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003. This is a three year pay for performance demonstration with physicians to promote the adoption and use of health information technology (HIT) to improve the quality of care for eligible chronically ill Medicare beneficiaries. MCMP targets small to medium sized primary care practices with up to 10 physicians. Practices must provide care to at least 50 Medicare beneficiaries. Physicians will receive payments for meeting or exceeding performance standards for quality of care. They will also receive an additional incentive payment for electronic submission of performance measures via their electronic health record (EHR) system. These payments are in addition to their normal payments for providing service to Medicare beneficiaries. The Office System Survey (OSS) will be used to assess progress of physician practices in implementation and use of EHRs and related HIT functionalities.

The EHR demonstration is authorized under section 402 of the Medicare Waiver Authority. The goal of this six year pay for performance demonstration is to foster the implementation and adoption of EHRs and HIT in order to improve the quality of care provided by physician practices. The EHRD expands upon the MCMP Demonstration and will test whether performance-based financial incentives (1) increase physician practices' adoption and use of electronic health records (EHRs), and (2) improve the quality of care that practices deliver to chronically ill patients. The EHRD targets small to medium sized primary care practices with up to 20 physicians. Practices must provide care to at least 50 Medicare beneficiaries. Approximately 2,400 practices will be enrolled in the demonstration across 12 sites. Practices will be randomly assigned to a treatment and control group. The OSS will be used to assess progress of physician practices in implementation and use of EHRs and related HIT functionalities, and to determine incentive payments for treatment practices. In-person and telephone discussions with community partners and physician practices will be used to learn about practices' experiences and

strategies in adopting and using EHRs, as well as the factors that help or hinder their efforts. *Form Number*: CMS–10273 (OMB# 0938—New); *Frequency*: Annually, Biennially and Once; *Affected Public*: Business or other forprofit; *Number of Respondents*: 3434; *Total Annual Responses*: 3434; *Total Annual Hours*: 2586.

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS' Web site at *http://www.cms.hhs.gov/Paperwork ReductionActof1995*, or E-mail your request, including your address, phone number, OMB number, and CMS document identifier, to *Paperwork@cms.hhs.gov*, or call the Reports Clearance Office on (410) 786– 1326.

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by *March 24, 2009:*

1. *Electronically*. You may submit your comments electronically to *http:// www.regulations.gov*. Follow the instructions for "Comment or Submission" or "More Search Options" to find the information collection document(s) accepting comments.

2. *By regular mail.* You may mail written comments to the following address: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number __, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

Dated: January 14, 2009.

Michelle Shortt,

Director, Regulations Development Group, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. E9–1435 Filed 1–22–09; 8:45 am] BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[CMS-2899-FN]

Medicare and Medicaid Programs; Approval of the Accreditation Commission for Health Care, Incorporated for Continued Deeming Authority for Home Health Agencies

AGENCY: Centers for Medicare & Medicaid Services (CMS), HHS.

ACTION: Final notice.

SUMMARY: This notice announces our decision to approve the Accreditation Commission for Health Care, Incorporated (ACHC) for continued recognition as a national accreditation program for home health agencies (HHAs) seeking to participate in the Medicare or Medicaid programs.

DATES: *Effective Date:* This final notice is effective February 24, 2009 through February 24, 2015.

FOR FURTHER INFORMATION CONTACT:

Lillian Williams, (410) 786–8636. Patricia Chmielewski, (410) 786–6899. SUPPLEMENTARY INFORMATION:

I. Background

Under the Medicare program, eligible beneficiaries may receive selected covered services from a home health agency (HHA) provided certain requirements are met. Sections 1861(m) and (o), 1891, and 1895 of the Social Security Act (the Act) authorize the Secretary to establish distinct criteria for facilities seeking designation as an HHA. Under this authority, the minimum requirements that an HHA must meet to participate in Medicare are set forth in regulations at 42 CFR part 484 and 42 CFR part 409, which determine the basis and scope of HHAcovered services, and the conditions for Medicare payment for home health care. Regulations concerning provider agreements are at 42 CFR part 489 and those pertaining to activities relating to the survey and certification of facilities are at 42 CFR part 488.

Generally, to enter into an agreement, an HHA must first be certified by a State survey agency as complying with conditions or requirements set forth in part 484 of our regulations. Then, the HHA is subject to regular surveys by a State survey agency to determine whether it continues to meet those requirements. There is an alternative, however, to surveys by State agencies.

Section 1865(a)(1) of the Act (as redesignated under section 125 of the Medicare Improvements for Patients and Providers Act of 2008 (MIPPA) (Pub. L. 110-275) provides that, if a provider entity demonstrates through accreditation by an approved national accreditation organization that all applicable Medicare conditions are met or exceeded, we may "deem" those provider entities as having met Medicare requirements. (We note that section 125 of MIPPA redesignated subsections (b) through (e) of subsection 1865 of the Act as (a) through (d) respectively.) Accreditation by an accreditation organization is voluntary